(FILE THEAPLOS SENTERED AT 12:48:41 ON 31 JUL 2002) - key terms (MORAXEL? OR M OR 1343 SEA FILE=HCAPLUS ABB=ON PLU=ON L1BRANHAMELL? OR M) (W) CATARRH? 56 SEA FILE=HCAPLUS ABB=ON PLU=ON L1 (5A) ANTIGEN L431 SEA FILE=HCAPLUS ABB=ON PLU=ON L4(S) VACCIN? r8 31 SEA FILE=HCAPLUS ABB=ON PLU=ON L8 AND (POLYPEPTIDE OR L9 PEPTIDE OR PROTEIN OR POLYPROTEIN) 22 SEA FILE=HCAPLUS ABB=ON PLU=ON L9 AND (ANTIBOD? OR Li2 T(W) (CELL OR LYMPHOCYT?)) L12 ANSWER 1 OF 22 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2001:255245 HCAPLUS DOCUMENT NUMBER: 134:265146 Cloning and characterization of outer membrane TITLE: protein OMP106 gene of Moraxella catarrhalis and its prophylactic, diagnostic and therapeutic uses Tucker, Kenneth; Plosila, Laura; Tillman, Ulrich INVENTOR(S): Antex Biologics Inc., USA PATENT ASSIGNEE(S): U.S., 49 pp., Cont.-in-part of U.S. Ser. No. SOURCE: 642,712. CODEN: USXXAM DOCUMENT TYPE: Patent LANGUAGE: English 2 FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	US 6214981	B1		US 1997-968685	19971112
	CN 1223549	Α	19990721	CN 1997-195990	19970428
	ZA 9703809	Α	19971201	ZA 1997-3809	19970502
				KR 1998-708845	
PRIO	RITY APPLN. INFO.	: .	US	1996-642712 A2	19960503
AB	The invention di	sclose	s the Moraxella	catarrhalis oute	r membrane
	protein-106 (OMP	106) p	olypeptide,		
	polypeptides der	ived t	herefrom (OMP10	6-derived	÷
	polypeptides), n	ucleot	ide sequences e	ncoding these	
	polypeptides, an	d anti	bodies that spe	cifically	
	bind the OMP106	polype	<pre>ptide and/or OM</pre>	P106-derived	
	polypeptides. A	lso di	sclosed are imm	unogenic, prophyl	actic
	or therapeutic c	ompns.	, including vac	cines, comprising	OMP106
	polypeptide and/	or OMP	106-derived pol	ypeptides.	
	The invention ad	dnl. d	iscloses method	s of inducing imm	une responses
	to M. catarrhali	s and	M. catarrhalis	OMP106 polypeptid	es
	and OMP106-deriv	ed pol	ypeptides in an	imals.	
REFE	RENCE COUNT:	21		1 CITED REFERENCE	
			FOR THIS RE	CORD. ALL CITATIO	NS AVAILABLE

IN THE RE FORMAT

HCAPLUS COPYRIGHT 2002 ACS

L12 ANSWER 2 OF 22 2001:168028 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 134:221433

Vaccine antigens of Moraxella TITLE:

Farn, Jacinta; Strugnell, Richard; Tennent, Jan INVENTOR(S): PATENT ASSIGNEE(S): Commonwealth Scientific and Industrial Research Organisation, Australia; The University of

> 308-4994 Shears Searcher :

Melbourne

SOURCE: PCT Int. Appl., 60 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PAT	TENT		KI	KIND DATE APPLICATION NO. DAT				DATE								
	WO	2001	0161	72	A1 20010308				WO 2000-AU1048 20000					0831			
		W:	AE,	AG,	AL,	AM,	AT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,
															GD,		_
			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	ΚZ,	LC,	LK,
•			LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,
			PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	ΤZ,
			UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZW,	AM,	AZ,	BY,	KG,	ΚZ,	MD,	RU,
			TJ,	TM													
		RW:	GH,	GM,	ΚE,	LS,	MW,	ΜZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,
			CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,
			BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG
	ΕP	1210	364		A.	1	2002	0605		E	P 20	00-9	5597	4	2000	0831	
		R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙT,	LI,	LU,	NL,	SE,	MC,
			PT,	ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL					
	BR	2000	0135	74	Α		2002	0611		Bl	R 20	00-1	3574		2000	0831	
PRIO	RITY	APP	LN.	INFO	. :					AU 19	999-	2571		Α	1999	0831	
	•		•						1	WO 20	000-	AU10	48	W	2000	0831	
AR	The	pre	sent	inve	entid	on r	elate	es to	o an	tiae	ns o	f Mo:	raxe.	lla,	in		

AB The present invention relates to antigens of Moraxella, in particular, Moraxella bovis, nucleic acid sequences encoding these antigens and formulations for use in raising an immune response against Moraxella.

REFERENCE COUNT:

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 3 OF 22 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:101183 HCAPLUS

DOCUMENT NUMBER:

134:161878

TITLE:

Moraxella catarrhalis BASB114 antigens and uses

thereof

INVENTOR(S):

Thonnard, Joelle

PATENT ASSIGNEE(S):

Smithkline Beecham Biologicals S.A., Belg.

SOURCE:

PCT Int. Appl., 82 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

1

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT	NO.		KI	ND :	DATE			A	PPLI	CATI	ои ис	ο.	DATE		
ETO 200	71 20010209			WO 2000-EP7293 20000727											
WO 200	TOORT	19	A	Ι.	2001	0200		W	0 20	UU-6.	P123.	3	2000	0121	
W:	AE,	AG,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,
	CN,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,
	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,
	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,
	UA,	ŪG,	US,	UZ,	VN,	YU,	ZA,	ZW,	AM,	AZ,	BY,	KG,	ΚZ,	MD,	RU,

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TJ, TM
           RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                             A1
                                   20020515
                                                       EP 2000-956338 20000727
      EP 1204678
           R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, MC, IE,
                 SI, LT, LV, FI, RO, MK, CY, AL
PRIORITY APPLN. INFO.:
                                                    GB 1999-17977
                                                                          A 19990730
                                                    WO 2000-EP7293
                                                                          W 20000727
      The invention provides BASB114 polypeptides and
AΒ
      polynucleotides encoding BASB114 polypeptides and methods
      for producing such polypeptides by recombinant techniques.
      Also provided are diagnostic, prophylactic and therapeutic uses.
                                        THERE ARE 1 CITED REFERENCES AVAILABLE FOR
REFERENCE COUNT:
                                1
                                        THIS RECORD. ALL CITATIONS AVAILABLE IN
                                         THE RE FORMAT
L12 ANSWER 4 OF 22 HCAPLUS COPYRIGHT 2002 ACS
                                2001:78537 HCAPLUS
ACCESSION NUMBER:
                                134:144470
DOCUMENT NUMBER:
                                A high molecular weight major outer membrane
TITLE:
                                protein of Moraxella and the gene
                                encoding it and the diagnosis, prophylaxis and
                                treatment of infection
                                Loosmore, Sheena M.; Sasaki, Ken; Yang,
INVENTOR(S):
                                Yan-Ping; Klein, Michel H.
                                Connaught Laboratories Limited, Can.
PATENT ASSIGNEE(S):
                                PCT Int. Appl., 247 pp.
SOURCE:
                                CODEN: PIXXD2
DOCUMENT TYPE:
                                Patent
                                English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                                        APPLICATION NO.
      PATENT NO.
                            KIND
                                    DATE
                                     -----
                                                        _____
                                                                              20000726
                                    20010201
                                                      WO 2000-CA870
      WO 2001007619
                            A1
           W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH,
                CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU,
                 TJ, TM
                GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
           RW: GH, GM,
                                                        EP 2000-951136 20000726
                                   20020508
      EP 1203082
                             A1
                AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL
                                                     US 1999-361619
                                                                           A2 19990727
PRIORITY APPLN. INFO.:
```

AB An isolated and purified outer membrane **protein** of a Moraxella strain, particularly M. catarrhalis, having a mol. mass of about 200 kDa, is provided by recombinant means. The about 200 kDa outer membrane **protein** as well as nucleic acid mols. encoding the same are useful in diagnostic applications and immunogenic compns., particularly for in vivo administration to a

Searcher: Shears 308-4994

WO 2000-CA870

W 20000726

host to confer protection against disease caused by a bacterial pathogen that produces the about 200 kDa outer membrane protein or produces a protein capable of inducing antibodies in a host specifically reactive with the about 200 kDa outer membrane protein. N-terminally and C-terminally truncated about 200 kDa proteins also are produced recombinantly. The gene was cloned from the 4223 strain by screening an expression library in .lambda.EMBL3 with antiserum to the protein. A series of overlapping fragments were obtained and assembled to give the full-length gene. The gene was then used as a probe to obtain the gene from a no. of different strains of the bacterium. Protein manufd. in Escherichia coli was obtained as inclusion bodies that could be resolubilized and used raise antiserum in mice and guinea pigs. The antiserum was bactericidal and could block the binding of the bacterium to animal cells. Comparison of the sequences of the G tract of genes from strains with different clumping activity indicated that the no. of G's in the tract affected levels of gene expression. Prepn. and characterization of N- and C-terminal truncation derivs. is described.

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR 3 THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

HCAPLUS COPYRIGHT 2002 ACS L12 ANSWER 5 OF 22

ACCESSION NUMBER: 2001:23521 HCAPLUS

DOCUMENT NUMBER: 135:194002

TITLE: Vaccines for Moraxella catarrhalis

McMichael, J. C. AUTHOR(S):

Wyeth-Lederle Vaccines, West Henrietta, NY, CORPORATE SOURCE:

14586-9728, USA

SOURCE: Vaccine (2000), 19(Suppl. 1), S101-S107

CODEN: VACCDE; ISSN: 0264-410X

PUBLISHER: Elsevier Science Ltd. Journal; General Review DOCUMENT TYPE:

LANGUAGE: English

A review with 53 refs. Vaccine development for M AB . catarrhalis is in the antigen identification stage. M. catarrhalis does not appear to synthesize secreted antigens such as exotoxins, nor does it appear to possess a carbohydrate capsule. Modified forms of these antigens are usually good vaccine components. There is some interest in whole bacterial cells and membrane fractions, but the search has largely focused on purified outer surface antigens. All of the present antigens have been selected based on the response seen in animals, although the antibody response seen in people exposed to the bacterium provides some guidance. The antibody response provides information related to the cross-strain preservation of epitopes and whether they are surface exposed. Antigens that elicit antibodies that have complement dependent bactericidal capacity, opsonophagocytic activity or interfere with one of the antigen's known functions such as adhesion or nutrient acquisition are particularly valued. In addn. to examg. the antibody response, some antigens have been evaluated in a murine pulmonary clearance model. Using these assays and model, several vaccine candidates have been identified. The antigens may be roughly classified by the function they serve the bacterium. One set appears to promote adhesion to host tissues and includes the

hemagglutinins, ubiquitous surface protein Al (UspAl), and possibly the CD protein. A second set is involved in nutrient acquisition. This set includes the lactoferrin binding protein A (LbpA) and lactoferrin binding protein B (LbpB), the transferrin binding protein A (TbpA) and transferrin binding protein B (TbpB), the CD and E porins, and the catarrhalis outer membrane protein B (CopB). A third set is comprised of antigens involved in virulence and it includes lipooligosaccharide (LOS) and the ubiquitous surface protein A2 (UspA2). Antigens of unknown function, such as the 200 K protein, may also be vaccine candidates.

REFERENCE COUNT:

53 THERE ARE 53 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 6 OF 22 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 2000:628168 HCAPLUS

DOCUMENT NUMBER: 133:221588

TITLE: Immunogenic compounds INVENTOR(S): Ruelle, Jean-louis

PATENT ASSIGNEE(S): Smithkline Beecham Biologicals S.A., Belg.

SOURCE: PCT Int. Appl., 97 pp.

CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE: Patent English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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KIND
                                                    DATE
          PATENT NO.
                                                                              APPLICATION NO.
                                                    _____
                                      · A1 20000908
                                                                           WO 2000-EP1468 20000223
          WO 2000052042
                 W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR,
                         CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU,
                 CU, CZ, DE, DK, DM, EE, ES, F1, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                          A1
                                                                              EP 2000-907603
                                                 20011219
                                                                                                            20000223
          EP 1163265
                 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,
                         PT, IE, FI
PRIORITY APPLN. INFO.:
                                                                          GB 1999-4559
                                                                                                        A 19990226
                                                                         WO 2000-EP1468
                                                                                                        W 20000223
```

AB The invention provides BASB081 polypeptides from Moraxella catarrhalis and polynucleotides encoding BASB081 polypeptides and methods for producing such

polypeptides by recombinant techniques. Also provided are

diagnostic, prophylactic and therapeutic uses.

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 7 OF 22 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2000:227773 HCAPLUS

DOCUMENT NUMBER: 132:250005

TITLE: Antigenic outer membrane protein OMP21

of Moraxella catarrhalis and the gene encoding it and their prophylactic, diagnostic and

therapeutic uses

INVENTOR(S):

Tucker, Kenneth; Tillmann, Ulrich F.

PATENT ASSIGNEE(S): SOURCE:

Antex Biologics Inc., USA PCT Int. Appl., 109 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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APPLICATION NO.
     PATENT NO.
                          KIND
                                  DATE
                                  -----
     WO 2000018910
                                  20000406
                                                  WO 1999-US22918 19991001
                          A1
              AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL,
               IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW,
               AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
          RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
               DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF,
               BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                                                        19991001
                                  20000417
                                                  AU 1999-64100
     AU 9964100
                           A1
     EP 1117779
                                  20010725
                                                    EP 1999-951716
                                                                        19991001
                           A1
              AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,
               PT, IE, SI, LT, LV, FI, RO
PRIORITY APPLN. INFO.:
                                                 US 1998-164714
                                                                      A 19981001
                                                 WO 1999-US22918 W 19991001
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The invention discloses the Moraxella catarrhalis outer membrane AΒ protein polypeptide and polypeptides derived therefrom (collectively "OMP21"), nucleotide sequences encoding said OMP21, and antibodies that specifically bind OMP21. Also disclosed are pharmaceutical compns. including prophylactic or therapeutic compns., which may be immunogenic compns. including vaccines, comprising OMP21, antibodies thereto or nucleotides encoding same. The invention addnl. discloses methods of inducing an immune response to M. catarrhalis and OMP21 in an animal, preferably a human, methods of treating and methods of diagnosing Moraxella infections in an animal, preferably a human, and kits therefor. The outer membrane proteins of several strains of M. catarrhalis were extd. with non-denaturing detergents (octyl glucoside or EmpigenBB.RTM.) and fractionated on SDS-polyacrylamide gels followed by transfer to PVDF membranes for N-terminal sequencing. The protein was antigenic in rabbits and conserved between strains of M. catarrhalis and related bacteria. Antisera to the protein mediated complement killing of M. catarrhalis. The gene, omp21, was cloned by PCR with degenerate primers and a knockout mutation created. The knockout strain showed weaker binding to cultured nasopharyngeal cells than did the wild type.

REFERENCE COUNT:

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 8 OF 22 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2000:191223 HCAPLUS

> 308-4994 Searcher Shears

132:233331 DOCUMENT NUMBER: Moraxella catarrhalis basb034 TITLE: polypeptides and utility in vaccine development and diagnosis Ruelle, Jean-louis INVENTOR(S): Smithkline Beecham Biologicals S.A., Belg. PATENT ASSIGNEE(S): PCT Int. Appl., 106 pp. SOURCE: CODEN: PIXXD2 DOCUMENT TYPE: Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: APPLICATION NO. KIND DATE DATE PATENT NO. -----WO 1999-EP6781 19990914 20000323 WO 2000015802 Α1 AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG AU 1999-58632 19990914 AU 9958632 A1 20000403 BR 1999-14492 19990914 20010626 BR 9914492 Α EP 1999-946171 19990914 20010711 EP 1114160 A1 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO NO 2001-1263 20010313 NO 2001001263 20010430 Α GB 1998-20002 19980914 Α PRIORITY APPLN. INFO.: WO 1999-EP6781 W 19990914 The invention provides BASB034 polypeptides and AB polynucleotides encoding BASB034 polypeptides and methods for producing such polypeptides by recombinant techniques. It is not uncommon to isolate Moraxella catarrhalis strains that are resistant to some or all of the std. antibiotics. The gene BASB034 was isolate from Moraxella catarrhalis strain ATCC43617 and other The non-coding flanking regions of the BASB034 gene were analyzed and exploited for modulation of BASB034 gene expression. Rflp patterns within this gene were found with the following restriction endonucleases: HphI, AluI, RsaI, EcoRV, and Sau3A1. vaccine is described comprising the gene BASB034 protein and at least one other Moraxella catarrhalis antigen. This may be used to generate an immune response. Antibodies specific for this antigen are discussed in the light of Moraxella catarrhalis infection detection and treatment and diagnosis. Also provided are diagnostic, prophylactic and therapeutic uses. THERE ARE 1 CITED REFERENCES AVAILABLE FOR REFERENCE COUNT: 1 THIS RECORD. ALL CITATIONS AVAILABLE IN

L12 ANSWER 9 OF 22 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2000:133833 HCAPLUS

DOCUMENT NUMBER: 132:176650

TITLE: Cloning of BASB023 antigen from Moraxella

THE RE FORMAT

catarrhalis

INVENTOR(S):

Thonnard, Joelle

PATENT ASSIGNEE(S):

Smithkline Beecham Biologicals S.A., Belg.

SOURCE:

PCT Int. Appl., 99 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

DATE APPLICATION NO. DATE PATENT NO. KIND _____ -----_____ 20000224 19990811 WO 2000009694 A1 WO 1999-EP5828 AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG AU 1999-54227 A1 20000306 19990811 AU 9954227 EP 1105492 A1 20010613 EP 1999-940192 19990811 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO

GB 1998-17824 A 19980814 PRIORITY APPLN. INFO.: WO 1999-EP5828 W 19990811

The invention provides BASB023 polypeptides and AB polynucleotides encoding BASB023 polypeptides from Moraxella catarrhalis (also named Branhamella catarrhalis) and methods for producing such polypeptides by recombinant techniques. BASB023 antigen is related by amino acid sequence homol, to Legionella adelaidensis macrophage infectivity potentiator polypeptide. Since Moraxella catarrhalis is responsible for several pathologies, the main ones being otitis media in infants and children and pneumonia in elderlies, the invention provides diagnostic, prophylactic and therapeutic uses for Moraxella infection.

REFERENCE COUNT:

2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 10 OF 22 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1999:736756 HCAPLUS

DOCUMENT NUMBER:

131:350252

TITLE:

SOURCE:

Moraxella catarrhalis antigenic proteins

and their use for immunization

INVENTOR(S):

Cripps, Allan William; Kyd, Jennelle

PATENT ASSIGNEE(S):

Cortecs (UK) Limited, UK PCT Int. Appl., 32 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.

KIND DATE APPLICATION NO. DATE

Searcher :

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308-4994

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A2
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     WO 9958563
                            19991229
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     EP 1077999
                       A2
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     NO 2000005670
                       Α
                                                            19980511
                                        GB 1998-10084
                                                         Α
PRIORITY APPLN. INFO.:
                                                            19990511
                                        WO 1999-GB1473
                                                         W
     Novel antigens of Branhamella
AB
     catarrhalis (also known as Moraxella catarrhalis) are
     provided, together with their use in vaccines as well as
     methods of diagnosis and/or detection. N-terminal and internal
    peptide sequences are provided for antigenic
    proteins of mol. mass 20, 30, 35, 44, and 71 kDa.
                      HCAPLUS COPYRIGHT 2002 ACS
L12 ANSWER 11 OF 22
                         1999:723176 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         131:347525
                         Moraxella catarrhalis Basb019 proteins
TITLE:
                         and genes from Moraxella catarrhalis and
                         antigens and antibodies and
                         therapeutic applications
INVENTOR(S):
                         Ruelle, Jean-Louis
                         SmithKline Beecham Biologicals S.A., Belg.
PATENT ASSIGNEE(S):
SOURCE:
                         PCT Int. Appl., 101 pp.
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
                         English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                            DATE
                                           APPLICATION NO.
                                                            DATE
     PATENT NO.
                      KIND
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     WO 9957277
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             SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW,
             AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE,
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CA 1999-2327316 19990503

CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

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AU 9939315
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     EP 1075521
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                        A2
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,
              PT, IE, FI
                                            GB 1998-9683
                                                              A 19980506
PRIORITY APPLN. INFO.:
                                                              W 19990503
                                            WO 1999-EP3038
     The invention provides Moraxella catarrhalis strain ATCC43617 gene
AB
     BASB019 polypeptides and polynucleotides encoding BASB019
     polypeptides and methods for producing such
     polypeptides by recombinant techniques. Variability within
     the BASB019 gene among several Moraxella catarrhalis strains was
     shown by RFLP anal. Also provided are diagnostic, prophylactic and
     therapeutic uses including prodn. of antisera to recombinant BASB019
     and vaccine prodn. and immunizations. A treatment of humans for
     Moraxella catarrhalis disease using antibody directed
     against Basb019 proteins is described. Lastly, screening
     assays for antagonists and agonists for BASB019 are described.
L12 ANSWER 12 OF 22 HCAPLUS COPYRIGHT 2002 ACS
                           1999:708913 HCAPLUS
ACCESSION NUMBER:
                           131:333042
DOCUMENT NUMBER:
                           Protein and DNA sequences of Moraxella
TITLE:
                           catarrhalis BASB011 gene, and uses thereof in
                           vaccine compositions and in assays for the
                           diagnosis of bacterial infections
                           Ruelle, Jean-louis
INVENTOR(S):
                           Smithkline Beecham Biologicals S.A., Belg.
PATENT ASSIGNEE(S):
                           PCT Int. Appl., 108 pp.
SOURCE:
                           CODEN: PIXXD2
DOCUMENT TYPE:
                           Patent
                           English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                       KIND DATE
                                               APPLICATION NO.
                                                                  DATE
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                       A1 · 19991104 WO 1999-EP2764 19990420
     WO 9955871
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              DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
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                        AA
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     AU 9940331
                         A1
                              19991116
                              20010131
                                               EP 1999-923457
                                                                  19990420
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A 19980423 W 19990420 WO 1999-EP2764 This invention provides the sequence of the Moraxella catarrhalis AΒ BASB011 gene, which encodes a protein that has homol. to the HtrA serine protease of Helicobacter pylori. The invention also relates to the use of an immunogenic fragment, preferably the extracellular domain, of the provided protein in a

EP 1071784

PRIORITY APPLN. INFO.:

A1

PT, IE, FI

AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,

GB 1998-8720

vaccine. The invention further relates to the use of the provided protein and/or gene in the diagnosis of bacterial

infections, esp. those of Moraxella.

REFERENCE COUNT:

THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN

THE RE FORMAT

HCAPLUS COPYRIGHT 2002 ACS L12 ANSWER 13 OF 22

ACCESSION NUMBER:

1999:554570 HCAPLUS

DOCUMENT NUMBER:

131:285063

TITLE:

Analysis of antigenic structure and human immune

response to outer membrane protein CD

of Moraxella catarrhalis

AUTHOR(S):

Murphy, Timothy F.; Kirkham, Charmaine;

DeNardin, Ernesto; Sethi, Sanjay

CORPORATE SOURCE:

Divisions of Infectious Diseases, Department of

Microbiology, State University of New York at

Buffalo, Buffalo, NY, 14215, USA

SOURCE:

Infection and Immunity (1999), 67(9), 4578-4585

CODEN: INFIBR; ISSN: 0019-9567

PUBLISHER:

American Society for Microbiology

DOCUMENT TYPE: LANGUAGE:

Journal English

Moraxella catarrhalis is an important cause of otitis media in AR children and lower respiratory tract infections in adults with chronic obstructive pulmonary disease (COPD). Outer membrane

protein CD (OMP CD) is a 45-kDa protein which is a potential vaccine antigen to prevent infections caused by M. catarrhalis. Eight monoclonal

antibodies were used to study the antigenic structure of the OMP CD mol. by assaying recombinant peptides corresponding to the sequence of the protein. This approach identified two surface-exposed epitopes, including one near the amino terminus (amino acids 25 to 44) and one in the central region of the mol. (amino acids 261 to 331). Assays with serum and sputum supernatants of adults with COPD revealed variable levels of antibodies to OMP CD among individuals. To det. which portions of the OMP CD mol. were recognized by human antibodies, three human serum samples were studied with six recombinant peptides which span the sequence of OMP CD. All three sera contained IgG antibodies which recognized exclusively the peptide corresponding to amino acids 203 to 260 by immunoblot assay. Adsorption expts. with whole bacteria established that some of the human antibodies are directed at surface-exposed epitopes on OMP CD. The authors conclude that OMP CD is a highly conserved mol. which contains at least two sep. epitopes which are exposed on the bacterial surface. While individual adults with COPD show variability in the immune response to OMP CD, a specific region of

REFERENCE COUNT:

the human immune response. THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE

IN THE RE FORMAT

the OMP CD mol. (amino acids 203 to 260) is important as a target of

HCAPLUS COPYRIGHT 2002 ACS L12 ANSWER 14 OF 22

43

ACCESSION NUMBER:

1999:83288 HCAPLUS

DOCUMENT NUMBER:

130:280494

TITLE:

Use of an isogenic mutant constructed in

Shears 308-4994 Searcher :

Moraxella catarrhalis to identify a protective

epitope of outer membrane protein B1 defined by monoclonal antibody 11C6

AUTHOR(S): Luke, Nicole R.; Russo, Thomas A.; Luther, Neal;

Campagnari, Anthony A.

CORPORATE SOURCE: Department of Microbiology, Center for Microbial

Pathogenesis, State University of New York at

Buffalo, Buffalo, NY, 14214, USA

SOURCE: Infection and Immunity (1999), 67(2), 681-687

CODEN: INFIBR; ISSN: 0019-9567 American Society for Microbiology

PUBLISHER: America:
DOCUMENT TYPE: Journal
LANGUAGE: English

Moraxella catarrhalis-induced otitis media continues to be a significant cause of infection in young children, prompting increased efforts at identifying effective vaccine antigens. authors have previously demonstrated that M. catarrhalis expresses specific outer membrane proteins (OMPs) in response to iron limitation and that this organism can utilize transferrin and lactoferrin for in vitro growth. One of these proteins, which binds human transferrin, is OMP B1. As the human host presents a naturally iron-limited environment, proteins, like OMP B1, which are expressed in response to this nutritional stress are potential vaccine antigens. In this study, the authors have developed monoclonal antibody (MAb) 11C6, which reacts to a surface-exposed epitope of OMP B1 expressed by M. catarrhalis 7169. This antibody was used to clone ompB1, and sequence anal. suggested that OMP B1 is the M. catarrhalis homolog to the transferrin binding protein B described for pathogenic Neisseriaceae, Haemophilus influenzae, Actinobacillus pleuropneumoniae, and M. catarrhalis. Expression of recombinant OMP B1 on the surface of Escherichia coli confers transferrin binding activity, confirming that this protein is likely involved in iron acquisition. In addn., ompB1 was used to construct an isogenic mutant in M. catarrhalis 7169. This mutant, termed 7169b12, was used as the control in bactericidal assays designed to det. if OMP B1 elicits protective antibodies. In the presence of MAb 11C6 and human complement, wild-type 7169 demonstrated a 99% decline in viability, whereas the ompB1 isogenic mutant was resistant to this bactericidal activity. Further anal. with MAb 11C6 revealed the presence of this OMP B1 epitope on 31% of the clin. isolates tested. These data suggest that OMP B1 is a

catarrhalis infections.
REFERENCE COUNT: 38

THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE

IN THE RE FORMAT

L12 ANSWER 15 OF 22 HCAPLUS COPYRIGHT 2002 ACS

potential vaccine antigen against M.

ACCESSION NUMBER:

1998:574816 HCAPLUS

DOCUMENT NUMBER:

129:313152

TITLE:

The transferrin binding protein B of

Moraxella catarrhalis elicits bactericidal antibodies and is a potential vaccine

antigen

AUTHOR (S):

Myers, Lisa E.; Yang, Yan-Ping; Du, Run-Pan; Wang, Qijun; Harkness, Robin E.; Schryvers, Anthony B.; Klein, Michel H.; Loosmore, Sheena

Μ.

CORPORATE SOURCE:

Pasteur Merieux Connaught Canada Research, North

York, ON, M2R 3T4, Can.

SOURCE:

Infection and Immunity (1998), 66(9), 4183-4192

CODEN: INFIBR; ISSN: 0019-9567 American Society for Microbiology

DOCUMENT TYPE:

Journal

PUBLISHER: LANGUAGE:

English

The transferrin binding protein genes (tbpA and tbpB) from AB two strains of Moraxella catarrhalis have been cloned and sequenced. The genomic organization of the M. catarrhalis transferrin binding protein genes is unique among known bacteria in that tbpA precedes tbpB and there is a third gene located between them. deduced sequences of the M. catarrhalis TbpA proteins from two strains were 98% identical, while those of the TbpB proteins from the same strains were 63% identical and 70% similar. The third gene, tentatively called orf3, encodes a protein of approx. 58 kDa that is 98% identical between the two strains. The tbpB genes from four addnl. strains of M. catarrhalis were cloned and sequenced, and two potential families of TbpB proteins were identified based on sequence similarities. Recombinant TbpA (rTbpA), rTbpB, and rORF3 proteins were expressed in Escherichia coli and purified. RTbpB was shown to retain its ability to bind human transferrin after transfer to a membrane, but neither rTbpA nor rORF3 did. Monospecific anti-rTbpA and anti-rTbpB antibodies were generated and used for immunoblot anal., which demonstrated that epitopes of M. catarrhalis TbpA and TbpB were antigenically conserved and that there was constitutive expression of the tbp genes. In the absence of an appropriate animal model, anti-rTbpA and anti-rTbpB antibodies were tested for their bactericidal activities. The anti-rTbpA antiserum was not bactericidal, but anti-rTbpB antisera were found to kill heterologous strains within the same family. Thus, if bactericidal ability is clin. relevant, a vaccine comprising multiple rTbpB antigens may protect against M.

L12 ANSWER 16 OF 22 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

catarrhalis disease.

1998:479556 HCAPLUS

DOCUMENT NUMBER:

129:108012

TITLE:

UspA1 and UspA2 antigens of Moraxella

catarrhalis

INVENTOR(S):

Hansen, Eric J.; Aebi, Christoph; Cope, Leslie

D.; Maciver, Isobel; Fiske, Michael J.;

Fredenburg, Ross

PATENT ASSIGNEE(S):

The Board of Regents, the University of Texas

System, USA

SOURCE:

PCT Int. Appl., 237 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. KIND DATE WO 9828333 A2 19980702 WO 1997-US23930 19971219

Searcher :

Shears

308-4994

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19990107
     WO 9828333
                         A3
             AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP,
             KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG,
              KZ, MD, RU, TJ, TM
          RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES,
              FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
              CI, CM,
                      GA, GN, ML, MR, NE, SN, TD, TG
                               19980717
                                               AU 1998-57201
                                                                  19971219
     AU 9857201
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                               20020502
                         B2
     AU 746442
                         A2
                               19991013
                                               EP 1997-953461
                                                                  19971219
     EP 948625
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              PT, IE, SI, LT, LV, FI, RO
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     CN 1251611
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     JP 2001515467
                                                                  19990615
     KR 2000057575
                         Α
                               20000925
                                               KR 1999-705332
                               20011030
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                                                                  19990621
     US 6310190
                         В1
                                            US 1996-33598P
                                                               Р
                                                                  19961220
PRIORITY APPLN. INFO.:
                                            WO 1997-US23930 W
                                                                  19971219
     The present invention discloses the existence of two novel
ΑB
     proteins UspA1 and UspA2, and their resp. genes uspA1 and
     uspA2. Each protein encompasses a region that is
     conserved between the two proteins and comprises an
     epitope that is recognized by MAb 17C7. One or more than one of
     these species may aggregate to form the very high mol. wt. form
     (i.e. greater than 200 kDa) of the UspA antigen. Compns. and both
     diagnostic and therapeutic methods for the treatment and study of M.
     catarrhalis are disclosed.
L12 ANSWER 17 OF 22 HCAPLUS COPYRIGHT 2002 ACS
                           1998:124040 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                           128:191575
TITLE:
                           Outer membrane protein B1 of Moraxella
                           catarrhalis
                           Campagnari, Anthony A.
INVENTOR(S):
                           Research Foundation of State University of New
PATENT. ASSIGNEE (S):
                           York, USA
                           PCT Int. Appl., 43 pp.
SOURCE:
                           CODEN: PIXXD2
DOCUMENT TYPE:
                            Patent
                           English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                                                APPLICATION NO.
                                                                  DATE
                        KIND
                               DATE
                               19980219
                                                WO 1997-US14596
                                                                  19970815
     WO 9806432
                         A1
          W: AU, CA, JP, MX
          RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
              PT, SE
                               19991221
                                                US 1996-698652
                                                                   19960816
     US .6004562
                         Α
                               19980306
                                                AU 1997-40757
                                                                  19970815
     AU 9740757
                         A1
PRIORITY APPLN. INFO .:
                                            US 1996-698652
                                                                  19960816
                                            WO 1997-US14596
                                                                  19970815
     An isolated and purified outer membrane protein B1, and
AB
     peptides formed therefrom, of Moraxella catarrhalis, are
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described. A method for the isolation and purifn. of outer membrane protein Bl from a bacterial strain that produces Bl protein, e.g. Moraxella catarrhalis, comprises growing the bacteria in culture in iron-depleted medium to enhance the expression of the Bl protein, harvesting the bacteria from the culture, extg. from the harvested bacteria a prepn. substantially comprising an outer membrane protein prepn., contacting the outer membrane prepn. with an affinity matrix contg. immobilized transferrin wherein Bl protein binds to the transferrin, and eluting the bound Bl protein from the transferrin. Disclosed are the uses of the Bl protein as an immunogen for vaccine formulations, and as antigens in diagnostic immunoassays.

L12 ANSWER 18 OF 22 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1997:596420 HCAPLUS

DOCUMENT NUMBER: 127:291797

TITLE: Antigenic heterogeneity and molecular analysis

of CopB of Moraxella (Branhamella) catarrhalis

AUTHOR(S): Sethi, S.; Surface, J. M.; Murphy, T. F.

CORPORATE SOURCE: Division of Pulmonary Medicine, State University

of New York at Buffalo, Buffalo, NY, USA

SOURCE: Infection and Immunity (1997), 65(9), 3666-3671

CODEN: INFIBR; ISSN: 0019-9567 American Society for Microbiology

PUBLISHER: American DOCUMENT TYPE: Journal

DOCUMENT TYPE: Journal LANGUAGE: English

Outer membrane protein (OMP) CopB, an iron-repressible AB 81-kDa major OMP of Moraxella (Branhamella) catarrhalis has been a major focus of investigation. To assess CopB as a potential vaccine antigen, the authors elucidated the degree of antigenic and sequence. heterogeneity in this protein among strains of M. catarrhalis. Two monoclonal antibodies, 1F5 and 2.9F, which bind to surface-exposed epitopes on CopB recognized 60 and 70% of the strains, resp. The degree of sequence heterogeneity in CopB was assessed by cloning and sequencing the CopB gene from two different strains of M. catarrhalis and comparing with the published sequence. There was 92 to 96% homol. between the sequences at the nucleotide level and 90 to 95% homol. at the amino acid level. variability in the protein sequence is confined mainly to three moderately variable regions. Restriction fragment length polymorphism (RFLP) anal. of the CopB genes obtained from 20 diverse strains by PCR was performed. Ninety percent of the potential restriction sites in the const. regions and 47% of the potential restriction sites in the variable regions were present in the 20 strains, indicating that the pattern of variable and const. areas in the CopB gene is a general pattern among strains of M. catarrhalis. The authors conclude that the CopB gene is largely conserved among strains of M. catarrhalis and contains discrete regions which show moderate heterogeneity among strains.

L12 ANSWER 19 OF 22 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1997:177696 HCAPLUS

DOCUMENT NUMBER: 126:249929

TITLE: The major outer membrane protein, CD, extracted from Moraxella (Branhamella)

catarrhalis is a potential
vaccine antigen that induces

bactericidal antibodies

Yang, Yan-ping; Myers, Lisa E.; McGuinness, AUTHOR(S): Ursula; Chong, Pele; Kwok, Yan; Klein, Michel

H.; Harkness, Robin E.

Research Center, Pasteur Merieux Connaught CORPORATE SOURCE:

Canada, 1755 Steeles Ave. West, North York, ON,

M2R 3T4, Can.

FEMS Immunology and Medical Microbiology (1997), SOURCE:

17(3), 187-199

CODEN: FIMIEV; ISSN: 0928-8244

Elsevier PUBLISHER: Journal DOCUMENT TYPE: LANGUAGE: English

The major outer membrane protein of Moraxella AB

(Branhamella) catarrhalis, CD, was detergent-extd. from the bacterial cell wall and purified to homogeneity in high yields by a simple process. The purified protein appeared to exhibit

immunogenic properties similar to those of native CD exposed on the surface of the bacterium. Antibodies to CD raised in mice specifically bound to intact B. catarrhalis, as detd. by flow cytometry anal. The IgG subclass distributions of anti-CD antibodies in sera from mice immunized with purified CD or with B. catarrhalis were also similar. CD was found to be

antigenically conserved among a panel of B. catarrhalis isolates, as demonstrated by the consistent reactivities of mouse anti-CD antisera with a common 60 kDa protein on immunoblots.

Furthermore, convalescent sera collected from patients with otitis media due to B. catarrhalis infection were found to be reactive with the CD protein by immunoblotting. Finally, the purified

protein induced antibodies in guinea pigs and mice

that exhibited in vitro bactericidal activity against the pathogen.

Therefore, the native CD outer membrane protein represents a potentially useful antigen for inclusion in a vaccine against B.

catarrhalis.

L12 ANSWER 20 OF 22 HCAPLUS COPYRIGHT 2002 ACS

1993:189964 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 118:189964

TITLE: Methods and compositions relating to useful

antigens of Moraxella catarrhalis

INVENTOR(S): Hansen, Eric J.; Helminen, Merja; Maciver,

Isobel

University of Texas System, USA PATENT ASSIGNEE(S):

SOURCE: PCT Int. Appl., 73 pp.

CODEN: PIXXD2.

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	ENT	NO.		KII	ND :	DATE			A.	PPLI	CATI	ои ис	э.	DATE		
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WO	9303	761		A.	L i	1993	0304		W	0 19	92-U	S686	9	1992	0814	
	W:	AT,	ΑU,	BB,	BG,	BR,	CA,	·CH,	CS,	DE,	DK,	ES,	FI,	GB,	HU,	JP,
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	RW:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	SE,
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US	5552	146		Α		1996	0903		U:	s 19	91-7	4559	1	1991	0815	

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19930316
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                             19940831
                       В1
                             19960724
     EP 612250
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     ES 2092696
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                             19961201
                                            ES 1992-918273
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                                            US 1993-25363
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                                                              19940214
                                            NO 1994-502
     NO 9400502
                       Α
                             19940328
     FI 9400681
                                            FI 1994-681
                       Α
                             19940407
                                                              19940214
                       Α
                             19980602
                                            US 1994-193150
                                                              19940919
     US 5759813
     US 5599693
                       Α
                             19970204
                                            US 1995-450002
                                                              19950525
                       Α
                             19991109
                                            US 1995-450351
                                                              19950525
     US 5981213
                       Α
                             20000509 .
                                            NO 2000-2413
                                                              20000509
     NO 2000002413
                                         US 1991-745591
                                                           A2 19910815
PRIORITY APPLN. INFO.:
                                         WO 1992-US6869
                                                           A 19920814
                                                           A3 19930302
                                         US 1993-25363
```

AB Selected antigenic proteins obtained from the outer membranes of M. catarrhalis are disclosed. These outer membrane proteins (OMPs) have mol. wts. of approx. 30 kDa, 80 kDa, and 200-700 kDa, resp. Studies demonstrated that monoclonal antibodies (MAbs) directed against these proteins confer a protective effect against infection by M. catarrhalis in animal models, demonstrating the potential usefulness of such antibodies in conferring passive immunity as well as the potential use of the OMPs (or variants thereof) in the prepn. of vaccines. DNA segments encoding the OMPs, methods for prepg. the antigens, and diagnostic methods are also disclosed. OMP isolation, anti-OMP MAb prodn., and cloning of genes for the OMPs are described.

L12 ANSWER 21 OF 22 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1

1993:17456 HCAPLUS

DOCUMENT NUMBER:

118:17456

TITLE:

Use of the purA gene as a selectable marker in

stabilization and integration of plasmid or

bacteriophage cloning vectors

INVENTOR(S):

Brey, Robert Newton, III; Fulginiti, James

Peter; Anilionis, Algis

PATENT ASSIGNEE(S):

American Cyanamid Co., USA

SOURCE:

Eur. Pat. Appl., 29 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

1

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 512260	A2	19921111	EP 1992-105887	19920406
EP 512260	A3	19930728		
R: AT, BE,	CH, DE	, DK, ES, FR,		, NL, PT, SE
AT 202800	E	20010715	AT 1992-105887	19920406
ES 2160573	Т3	20011116	ES 1992-105887	19920406
JP 05192161	A2	19930803	JP 1992-134375	19920428
NO 9201729	Α	19921104	NO 1992-1729	19920430
CA 2067862	AA	19921104	CA 1992-2067862	19920501

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19920501
                       A1
                             19921105
                                            AU 1992-15959
    AU 9215959
     AU 654347
                             19941103
                       В2
                             19990706
                                            US 1995-380297
                                                              19950130
     US 5919663
                       Α
                             19991005
                                            US 1995-448907
                                                              19950524
     US 5961983
                       Α
PRIORITY APPLN. INFO.:
                                         US 1991-695706
                                                          Α
                                                              19910503
                                         US 1994-204903
                                                           B1 19940302
                                         US 1995-380297
                                                           A3 19950130
```

AB Host bacteria carrying deletions in the purA gene (for adenylosuccinate synthetase) are used as hosts for cloning vectors carrying the purA gene as a selectable marker. The vector is stabilized by selection and the purA gene also acts as a site for integration of the plasmid. The use of these vectors does not involve the use of antibiotic resistance markers and is therefore particularly suitable for hosts used in live vaccines. A pUC8-based plasmid carrying the Escherichia coli purA gene and the gene for the nontoxic subunit of the E. coli heat-labile enterotoxin was constructed and introduced into Salmonella dublin, S. typhimurium or Salmonella vaccine strains carrying deletions in the purA gene and transformants selected on minimal medium. This plasmid was maintained in cultures grown on a minimal medium without loss for 80 generations but lost rapidly in the absence of selection (1% retention in 40 generations). When the purA gene was used in integrating vectors the prototrophic phenotype was 100% stable for at least 80 generations in the presence or absence of selection.

L12 ANSWER 22 OF 22 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1990:510481 HCAPLUS

DOCUMENT NUMBER:

113:110481

TITLE:

Fusion proteins of flagellin and

heterologous epitopes and attenuated bacteria

expressing the chimeric genes as vaccines

Marjarian, William Robert; Stocker, Bruce Arnold INVENTOR(S):

Dunbar; Newton, Salete Maria Cardozo

Praxis Biologics, Inc., USA; Leland Stanford

PATENT ASSIGNEE(S): Junior University

PCT Int. Appl., 137 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	TENT NO.		KIND	DATE		APPLICATION NO.	DATE
WO	8910967			19891116		WO 1989-US1932	19890505
	W: AU,		FI, JP	•			
	RW: AT,	BE,	CH, DE	, FR, GB,	IT,	LU, NL, SE	
ΑU	8936979		Al	19891129		AU 1989-36979	19890505
ΑU	637049		B2	19930520			
ΕP	419513		A1	19910403		EP 1989-906507	19890505
ΕP	419513		B1	19950426			
	R: AT,	BE,	CH, DE	, FR, GB,	IT,	LI, NL, SE	
JΡ	04502402		Т2	19920507		JP 1989-505981	19890505
JΡ	2793673	-	B2	19980903			
ΑТ	121782		E	19950515		AT 1989-906507	19890505
DK	9002633		Α	19910104		DK 1990-2633	19901102
NO	9004806		Α	19910103		NO 1990-4806	19901105
US	6130082		A	20001010		US 1992-837668	19920214

PRIORITY APPLN. INFO.:

US 1988-190570 A 19880505

US 1989-348430 B1 19890505

WO 1989-US1932 A 19890505

AB Fusion proteins of flagellin and an antigenic epitope prepd. by expression of the chimeric gene are used as vaccines. Similarly, the bacterium expressing the chimeric gene is also used in vaccines. Vertebrate hosts can be immunized by administering an invasive, but attenuated, bacterium that is transfected with a recombinant DNA encoding the fusion protein to elicit cellular or humoral immune response. Expression of heterologous parasitic, bacterial, and viral epitopes, e.g.malarial circumsporozoite protein antigen, the B subunit of cholera toxin, the epitope of the CRM197 protein (residues 366-383; a mutant or Diptheria toxin) hepatitis B virus surface antigen, and rotavirus VP7 antigen, with Salmonella flagellin in attenuated Salmonella were demonstrated and their immunogenicity obsd.

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(FILE OMEDLINE, BIOSIS, EMBASE, WPIDS, CONFSCI, SCISEARCH,
     JICST-EPIUS, JAPIO' ENTERED AT 12:49:48 ON 31 JUL 2002)
           1343 SEA FILE=HCAPLUS ABB=ON PLU=ON
                                                  (MORAXEL? OR M OR
L1
                BRANHAMELL? OR M) (W) CATARRH?
            · 56 SEA FILE=HCAPLUS ABB=ON PLU=ON
                                                 L1 (5A) ANTIGEN
L4
L8
             31 SEA FILE=HCAPLUS ABB=ON PLU=ON
                                                  L4(S) VACCIN?
             31 SEA FILE=HCAPLUS ABB=ON PLU=ON L8 AND (POLYPEPTIDE OR
L9
                PEPTIDE OR PROTEIN OR POLYPROTEIN)
             70 SEA L9
L10
             40 DUP REM L10 (30 DUPLICATES REMOVED)
L11
             37 SEA L11 AND (ANTIBOD? OR T(W) (CELL OR LYMPHOCYT?))
L13
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L13 ANSWER 1 OF 37 MEDLINE

ACCESSION NUMBER: 2001381129 MEDLINE

DOCUMENT NUMBER: 21108937 PubMed ID: 11163472 TITLE: Vaccines for Moraxella catarrhalis.

AUTHOR: McMichael J C

.CORPORATE SOURCE: Wyeth-Lederle Vaccines, 211 Bailey Road, West

Henrietta, NY 14586-9728, USA.. mcmichj@war.wyeth.com

SOURCE: VACCINE, (2000 Dec 8) 19 Suppl 1 S101-7. Ref: 53

Journal code: 8406899. ISSN: 0264-410X.

Doubled Code: 0400000 155N. 0204 4.

PUB. COUNTRY: England: United Kingdom

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

(REVIEW, TUTORIAL)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200107

ENTRY DATE: Entered STN: 20010709

Last Updated on STN: 20010709 Entered Medline: 20010705

AB Vaccine development for Moraxella

catarrhalis is in the antigen identification

stage. M. catarrhalis does not appear to synthesize secreted antigens such as exotoxins, nor does it appear to possess a carbohydrate capsule. Modified forms of these antigens are usually good vaccine components. There is some interest in whole bacterial cells and membrane fractions, but the

interest in whole bacterial cells and membrane fractions, but the search has largely focused on purified outer surface antigens. All of the present antigens have been selected based on the response

seen in animals, although the antibody response seen in people exposed to the bacterium provides some guidance. The antibody response provides information related to the cross-strain preservation of epitopes and whether they are surface exposed. Antigens that elicit antibodies that have complement dependent bactericidal capacity, opsonophagocytic activity or interfere with one of the antigen's known functions such as adhesion or nutrient acquisition are particularly valued. In addition to examining the antibody response, some antigens have been evaluated in a murine pulmonary clearance model. Using these assays and model, several vaccine candidates have been identified. The antigens may be roughly classified by the function they serve the bacterium. One set appears to promote adhesion to host tissues and includes the hemagglutinins, ubiquitous surface protein Al (UspAl), and possibly the CD protein. A second set is involved in nutrient acquisition. This set includes the lactoferrin binding protein A (LbpA) and lactoferrin binding protein B (LbpB), the transferrin binding protein A (TbpA) and transferrin binding protein B (TbpB), the CD and E porins, and the Catarrhalis outer membrane protein B (CopB). A third set is comprised of antigens involved in virulence and it includes lipooligosaccharide (LOS) and the ubiquitous surface protein A2 (UspA2). Antigens of unknown function, such as the 200K protein, may also be vaccine candidates. The antigens that are most suitable will be determined in clinical studies that are only beginning now.

L13 ANSWER 2 OF 37 MEDLINE

ACCESSION NUMBER: 2000036213 MEDLINE

DOCUMENT NUMBER: 20036213 PubMed ID: 10571435

TITLE: Antibody response to outer membrane

proteins of Moraxella catarrhalis in children

with otitis media.

AUTHOR: Mathers K; Leinonen M; Goldblatt D

CORPORATE SOURCE: Immunobiology Unit, Institute of Child Health,

London, UK.

SOURCE: PEDIATRIC INFECTIOUS DISEASE JOURNAL, (1999 Nov) 18

(11) 982-8.

Journal code: 8701858. ISSN: 0891-3668.

PUB. COUNTRY: United States
DOCUMENT TYPE: (CLINICAL TRIAL)

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199912

ENTRY DATE: Entered STN: 20000113

Last Updated on STN: 20000113 Entered Medline: 19991203

AB BACKGROUND: Moraxella catarrhalis is an important cause of bacterial otitis media, and a vaccine to prevent this disease would be highly desirable. Analysis of the dominant antigens on the surface of M. catarrhalis recognized by the human immune response to infection might aid in such a search. Such analysis would be most informative when studied in the eventual target age group for the vaccine; thus we have studied the immune response to M. catarrhalis in infants with otitis media. METHODS: Eighteen infants (mean age, 9.4 months) experiencing an

episode of otitis media caused by M. catarrhalis were studied. Acute and convalescent antibody responses were studied by whole cell enzyme-linked immunosorbent assay (heterologous strain) and by immunoblotting of outer membrane proteins (OMPs). RESULTS:

Specific IgG was detected in 17% of acute serum samples and in 61% of convalescent sera. A rise in specific IgG was detected in 10 of 12 (83%) children 8 months of age or older, compared with 1 of 6 (17%) in younger patients (P = 0.0128). Immunoblotting revealed antibody binding to several OMPs with some detectable cross-reactivity. Four dominant OMP targets were identified, corresponding to UspA, TbpB, CopB and a approximately 60-kDa protein. CONCLUSIONS: A combination of antigens might form the most suitable basis for a M. catarrhalis vaccine designed to prevent otitis media in this age group.

L13 ANSWER 3 OF 37 MEDLINE

ACCESSION NUMBER: 1999386849 MEDLINE

DOCUMENT NUMBER: 99386849 PubMed ID: 10456903

TITLE: Analysis of antigenic structure and human immune

response to outer membrane protein CD of

Moraxella catarrhalis.

AUTHOR: Murphy T F; Kirkham C; DeNardin E; Sethi S

CORPORATE SOURCE: Divisions of Infectious Diseases, School of Medicine

and Biomedical Sciences, State University of New York

at Buffalo, Buffalo, New York 14215, USA..

murphyt@acsu.buffalo.edu

CONTRACT NUMBER: AI28304 (NIAID)

SOURCE: INFECTION AND IMMUNITY, (1999 Sep) 67 (9) 4578-85.

Journal code: 0246127. ISSN: 0019-9567.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199910

ENTRY DATE: Entered STN: 19991014

Last Updated on STN: 19991014 Entered Medline: 19991005

Moraxella catarrhalis is an important cause of otitis media in AΒ children and lower respiratory tract infections in adults with chronic obstructive pulmonary disease (COPD). Outer membrane protein CD (OMP CD) is a 45-kDa protein which is a potential vaccine antigen to prevent infections caused by M. catarrhalis. Eight monoclonal antibodies were used to study the antigenic structure of the OMP CD molecule by assaying recombinant peptides corresponding to the sequence of the protein. This approach identified two surface-exposed epitopes, including one near the amino terminus (amino acids 25 to 44) and one in the central region of the molecule (amino acids 261 to 331). Assays with serum and sputum supernatants of adults with COPD revealed variable levels of antibodies to OMP CD among individuals. To determine which portions of the OMP CD molecule were recognized by human antibodies, three human serum samples were studied with six recombinant peptides which span the sequence of OMP CD. All three sera contained immunoglobulin G antibodies which recognized exclusively the peptide corresponding to amino

acids 203 to 260 by immunoblot assay. Adsorption experiments with whole bacteria established that some of the human **antibodies**

are directed at surface-exposed epitopes on OMP CD. We conclude that OMP CD is a highly conserved molecule which contains at least two separate epitopes which are exposed on the bacterial surface. While individual adults with COPD show variability in the immune response to OMP CD, a specific region of the OMP CD molecule (amino acids 203 to 260) is important as a target of the human immune response.

L13 ANSWER 4 OF 37 MEDLINE

ACCESSION NUMBER: 1999115543 MEDLINE

DOCUMENT NUMBER: 99115543 PubMed ID: 9916077

TITLE: Use of an isogenic mutant constructed in Moraxella

catarrhalis To identify a protective epitope of outer

membrane protein B1 defined by monoclonal

antibody 11C6.

AUTHOR: Luke N R; Russo T A; Luther N; Campagnari A A

CORPORATE SOURCE: Department of Microbiology, State University of New

York at Buffalo, Buffalo, New York 14214, USA.

INFECTION AND IMMUNITY, (1999 Feb) 67 (2) 681-7.

Journal code: 0246127. ISSN: 0019-9567.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

SOURCE:

FILE SEGMENT: Priority Journals OTHER SOURCE: GENBANK-AF105251

ENTRY MONTH: 199903

ENTRY DATE: Entered STN: 19990324

Last Updated on STN: 19990324 Entered Medline: 19990309

Moraxella catarrhalis-induced otitis media continues to be a AB significant cause of infection in young children, prompting increased efforts at identifying effective vaccine antigens. We have previously demonstrated that M. catarrhalis expresses specific outer membrane proteins (OMPs) in response to iron limitation and that this organism can utilize transferrin and lactoferrin for in vitro growth. One of these proteins, which binds human transferrin, is OMP B1. As the human host presents a naturally iron-limited environment, proteins, like OMP B1, which are expressed in response to this nutritional stress are potential vaccine antigens. In this study, we have developed monoclonal antibody (MAb) 11C6, which reacts to a surface-exposed epitope of OMP B1 expressed by M. catarrhalis 7169. This antibody was used to clone ompB1, and sequence analysis suggested that OMP B1 is the M. catarrhalis homologue to the transferrin binding protein B described for pathogenic Neisseriaceae, Haemophilus influenzae, Actinobacillus pleuropneumoniae, and M. catarrhalis. Expression of recombinant OMP B1 on the surface of Escherichia coli confers transferrin binding activity, confirming that this protein is likely involved in iron acquisition. In addition, ompB1 was used to construct an isogenic mutant in M. catarrhalis 7169. This mutant, termed 7169b12, was used as the control in bactericidal assays designed to determine if OMP B1 elicits protective antibodies. In the presence of MAb 11C6 and human complement, wild-type 7169 demonstrated a 99% decline in viability, whereas the ompB1 isogenic mutant was resistant to this bactericidal activity. Further analysis with MAb 11C6 revealed the presence of this OMP B1 epitope on 31% of the clinical isolates tested. These data suggest that OMP B1 is a

potential vaccine antigen against M. catarrhalis infections.

L13 ANSWER 5 OF 37 MEDLINE

ACCESSION NUMBER: 1998380363 MEDLINE

DOCUMENT NUMBER: 98380363 PubMed ID: 9712766

TITLE: The transferrin binding protein B of

Moraxella catarrhalis elicits bactericidal

antibodies and is a potential vaccine

antigen.

AUTHOR: Myers L E; Yang Y P; Du R P; Wang Q; Harkness R E;

Schryvers A B; Klein M H; Loosmore S M

CORPORATE SOURCE: Pasteur Merieux Connaught Canada Research, North

York, Ontario, Canada M2R 3T4.

SOURCE: INFECTION AND IMMUNITY, (1998 Sep) 66 (9) 4183-92.

Journal code: 0246127. ISSN: 0019-9567.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

OTHER SOURCE: GENBANK-AF039311; GENBANK-AF039312; GENBANK-AF039313;

GENBANK-AF039314; GENBANK-AF039315; GENBANK-AF039316

ENTRY MONTH: 199810

ENTRY DATE: Entered STN: 19981020

Last Updated on STN: 19981020

Entered Medline: 19981002

The transferrin binding protein genes (tbpA and tbpB) from AB two strains of Moraxella catarrhalis have been cloned and sequenced. The genomic organization of the M. catarrhalis transferrin binding protein genes is unique among known bacteria in that tbpA precedes tbpB and there is a third gene located between them. The deduced sequences of the M. catarrhalis TbpA proteins from two strains were 98% identical, while those of the TbpB proteins from the same strains were 63% identical and 70% similar. The third gene, tentatively called orf3, encodes a protein of approximately 58 kDa that is 98% identical
between the two strains. The tbpB genes from four additional strains of M. catarrhalis were cloned and sequenced, and two potential families of TbpB proteins were identified based on sequence similarities. Recombinant TbpA (rTbpA), rTbpB, and rORF3 proteins were expressed in Escherichia coli and purified. rTbpB was shown to retain its ability to bind human transferrin after transfer to a membrane, but neither rTbpA nor rORF3 did. Monospecific anti-rTbpA and anti-rTbpB antibodies were generated and used for immunoblot analysis, which demonstrated that epitopes of M. catarrhalis TbpA and TbpB were antigenically conserved and that there was constitutive expression of the tbp genes. In the absence of an appropriate animal model, anti-rTbpA and anti-rTbpB antibodies were tested for their bactericidal activities. The anti-rTbpA antiserum was not bactericidal, but anti-rTbpB antisera were found to kill heterologous strains within the same family. Thus, if bactericidal ability is clinically relevant, a vaccine comprising multiple rTbpB antigens may protect against M. catarrhalis disease.

L13 ANSWER 6 OF 37 MEDLINE

ACCESSION NUMBER: 97296466 MEDLINE

DOCUMENT NUMBER: 97296466 PubMed ID: 9152030

TITLE: Moraxella (Branhamella) catarrhalis--clinical and

molecular aspects of a rediscovered pathogen.

AUTHOR: Enright M C; McKenzie H

CORPORATE SOURCE: Department of Biological Sciences, University of

Sussex, Falmer, Brighton.

SOURCE: JOURNAL OF MEDICAL MICROBIOLOGY, (1997 May) 46 (5)

360-71. Ref: 129

Journal code: 0224131. ISSN: 0022-2615.

PUB. COUNTRY: ENGLAND: United Kingdom

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

(REVIEW, TUTORIAL)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199705

ENTRY DATE: Entered STN: 19970609

Last Updated on STN: 19970609 Entered Medline: 19970529

Since its discovery at the end of the nineteenth century, Moraxella AB (Branhamella) catarrhalis has undergone several changes of nomenclature and periodic changes in its perceived status as either a commensal or a pathogen. Molecular analysis based on DNA hybridisation or 16S rDNA sequence comparisons has established its phylogenetic position as a member of the Moraxellaceae and shown that it is related more closely to Acinetobacter spp. than to the genus Neisseria in which it was placed formerly. However, confusion with phenotypically similar Neisseria spp. can occur in the routine diagnostic laboratory if appropriate identification tests are not performed. M. catarrhalis is now accepted as the third commonest pathogen of the respiratory tract after Streptococcus pneumoniae and Haemophilus influenzae. It is a significant cause of otitis media and sinusitis in children and of lower respiratory tract infections in adults, especially those with underlying chest disease. Nosocomial spread of infection, especially within respiratory wards, has been reported. Invasive infection is uncommon, but analysis of reports for England and Wales between 1992 and 1995 revealed 89 cases of M. catarrhalis bacteraemia, with the peak incidence in children aged 1-2 years. Carriage rates of M. catarrhalis are high in children and in the elderly, but its role as a commensal organism has probably been overstated in the past. Approximately 90% of strains are now beta-lactamase positive and, given that the first such strain was reported in 1976, this represents a dramatic increase in frequency over the last 20 years which has not been paralleled in any other species. The BRO-1 and BRO-2 beta-lactamase enzymes of M. catarrhalis are found in other Moraxellaceae, but are not related to beta-lactamases of any other species and their origin is therefore unknown. Molecular and typing studies have shown that the M. catarrhalis species is genetically heterogeneous and these methods have aided epidemiological investigation. Studies of factors that may be related to pathogenicity have shown the existence of three serotypes of lipooligosaccharide and the presence of fimbriae and a possible capsule. Some strains are serum-resistant, probably by virtue of interference with complement action, whilst transferrin- and lactoferrin-binding proteins enable the organism to obtain iron from its environment. An antibody response in humans to various M. catarrhalis antigens, including highly conserved outer-membrane

proteins, has been demonstrated. Increased understanding of the organism's pathogenic properties and the host response to it may help to identify suitable vaccine targets or lead to other strategies to prevent infection. Whilst it remains, at present, the third most important respiratory pathogen, the impact of immunisation strategies for other organisms may change this position. The speed with which M. catarrhalis acquired beta-lactamase demonstrates the capacity of this organism to surprise us.

L13 ANSWER 7 OF 37 MEDLINE

ACCESSION NUMBER:

97247713 MEDLINE

DOCUMENT NUMBER:

97247713 PubMed ID: 9093840

TITLE:

The major outer membrane protein, CD, extracted from Moraxella (Branhamella) catarrhalis is a potential vaccine antigen that induces bactericidal

antibodies.

AUTHOR:

Yang Y P; Myers L E; McGuinness U; Chong P; Kwok Y;

Klein M H; Harkness R E

CORPORATE SOURCE:

Research Center, Pasteur Merieux Connaught Canada, North York, Ont., Canada.. ypyang@ca.pmc-vacc.com

SOURCE:

FEMS IMMUNOLOGY AND MEDICAL MICROBIOLOGY, (1997 Mar)

17 (3) 187-99.

Journal code: 9315554. ISSN: 0928-8244.

PUB. COUNTRY:

Netherlands

DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

199705

ENTRY DATE:

Entered STN: 19970609

Last Updated on STN: 19970609 Entered Medline: 19970529

The major outer membrane protein of Moraxella AΒ (Branhamella) catarrhalis, CD, was detergent-extracted from the bacterial cell wall and purified to homogeneity in high yields by a simple process. The purified protein appeared to exhibit immunogenic properties similar to those of native CD exposed on the surface of the bacterium. Antibodies to CD raised in mice specifically bound to intact B. catarrhalis, as determined by flow cytometry analysis. The IgG subclass distributions of anti-CD antibodies in sera from mice immunized with purified CD or with B. catarrhalis were also similar. CD was found to be antigenically conserved among a panel of B. catarrhalis isolates, as demonstrated by the consistent reactivities of mouse anti-CD antisera with a common 60 kDa protein on immunoblots. Furthermore, convalescent sera collected from patients with otitis media due to B. catarrhalis infection were found to be reactive with the CD protein by immunoblotting. Finally, the purified protein induced antibodies in guinea pigs and mice that exhibited in vitro bactericidal activity against the pathogen. Therefore, the native CD outer membrane protein represents a potentially useful antigen for inclusion in a vaccine against B. catarrhalis.

L13 ANSWER 8 OF 37

MEDLINE

ACCESSION NUMBER:

93329207 MEDLINE

DOCUMENT NUMBER:

93329207 PubMed ID: 8335988

Effect of immunization of pulmonary clearance of TITLE:

Moraxella catarrhalis in an animal model.

Maciver I; Unhanand M; McCracken G H Jr; Hansen E J AUTHOR:

Dept. of Microbiology, University of Texas CORPORATE SOURCE:

Southwestern Medical Center, Dallas 75235-9048. JOURNAL OF INFECTIOUS DISEASES, (1993 Aug) 168 (2) SOURCE:

469-72.

Journal code: 0413675. ISSN: 0022-1899.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE) DOCUMENT TYPE:

LANGUAGE: English

Abridged Index Medicus Journals; Priority Journals FILE SEGMENT:

ENTRY MONTH: 199308

Entered STN: 19930903 ENTRY DATE:

Last Updated on STN: 19970203 Entered Medline: 19930824

A murine model for pulmonary clearance of Moraxella catarrhalis was AB used to determine whether immunization could enhance clearance of this organism from the lungs. Animals actively immunized with outer membrane vesicles of M. catarrhalis cleared an endobronchial challenge with the homologous strain more quickly than did sham-immunized control animals. Western blot analysis of both this immune mouse serum and rabbit antiserum raised against outer membrane vesicles of M. catarrhalis indicated that antibodies were present to both outer membrane protein and lipooligosaccharide antigens. Passive immunization of mice with the immune rabbit serum resulted in enhanced pulmonary clearance of both homologous and heterologous strains of M. catarrhalis, indicating the involvement of serum antibody in this clearance process and the existence of conserved surface antigens in the two different M . catarrhalis strains. These results suggest that this model system may be useful for the identification of vaccine candidates among the surface antigens of M.

L13 ANSWER 9 OF 37 BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC.

ACCESSION NUMBER: 2002:201461 BIOSIS PREV200200201461 DOCUMENT NUMBER:

catarrhalis.

Intranasal immunization with detoxified TITLE:

lipooligosaccharides from Moraxella catarrhalis

conjugated to a protein elicit protection

in a mouse model of colonization.

Jiao, X. (1); Hirano, T. (1); Hou, Y. (1); Gu, X. (1) AUTHOR(S):

(1) Laboratory of Immunology, National Institute on CORPORATE SOURCE: Deafness and Other Communication Disorders, National

Institutes of Health, Rockville, MD USA

Abstracts of the General Meeting of the American SOURCE:

Society for Microbiology, (2001) Vol. 101, pp. 302. http://www.asmusa.org/mtgsrc/generalmeeting.htm.

Meeting Info.: 101st General Meeting of the American Society for Microbiology Orlando, FL, USA May 20-24,

2001

ISSN: 1060-2011.

DOCUMENT TYPE:

Conference

LANGUAGE:

English

Moraxella catarrhalis is a significant cause of otitis media in

Searcher : 308-4994 Shears

children. Lipooligosaccharide (LOS) is a major surface antigen of M. catarrhalis and a potential vaccine candidate. But little is known about the mucosal immune responses of detoxified LOS (dLOS)-protein conjugate vaccines and their potential roles on mucosal surfaces. In order to address these issues, BALB/c mice were immunized intranasally with a mixture of dLOS-CRM (the diphtheria toxin cross-reactive mutant protein) and cholera toxin (CT) as an adjuvant, dLOS plus CT, or CT only. After immunization, the animals were aerosolly challenged with M. catarrhalis strain 25238. Immunization with dLOS-CRM generated a significant increase in secreting IgA and IgG in nasal washes, bronchoalveolar lavage and saliva, and serum IgG, IgM and IgA against LOS of M. catarrhalis as detected by an enzyme-linked immunosorbent assay (ELISA). The dLOS-CRM elicited LOS-specific IgA, IgG, and IgM antibody -forming cells (AFCs) in different lymphoid tissues as measured by an enzyme-linked immunospot (ELISPOT) assay. LOS-specific IgA AFCs were found in the nasal passages, spleens, nasal-associated lymphoid tissues (NALT), cervical lymph nodes (CLN), lungs, and small intestines. LOS-specific IgG and IgM AFCs were only detected in the spleens, CLN, and nasal passages. Furthermore, the dLOS-CRM vaccine generated an 80% bacterial clearance in the nasopharynx and lungs when compared to the controls (P<0.01) following an aerosol challenge with the homologous strain 25238. Intriguingly, intranasal immunization with dLOS-CRM containing CT showed a higher level of bacterial clearance in both sites when compared to subcutaneous injections with dLOS-CRM plus a Ribi adjuvant. These data indicate that dLOS-CRM induces specific mucosal and systemic immunity against M. catarrhalis through intranasal immunization, and provides effective bacterial clearance in the mouse nasopharynx and lungs. Therefore, this may be an efficient route for vaccines to prevent otitis media and lower respiratory tract infections caused by M. catarrhalis.

L13 ANSWER 10 OF 37 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V.

ACCESSION NUMBER: 2002023188 EMBASE

TITLE: Moraxella catarrhalis: From emerging to established

pathogen.

AUTHOR: Verduin C.M.; Hol C.; Fleer A.; Van Dijk H.; Van

Belkum A.

CORPORATE SOURCE: C.M. Verduin, Department of Medical Microbiology,

Erasmus University Medical Center, Rotterdam EMCR,

Dr. Molewaterplein 40, 3015 GD Rotterdam,

Netherlands. verduin@bacl.azr.nl

SOURCE: Clinical Microbiology Reviews, (2002) 15/1 (125-144).

Refs: 256

ISSN: 0893-8512 CODEN: CMIREX

COUNTRY: United States

DOCUMENT TYPE: Journal; General Review FILE SEGMENT: 004 Microbiology

026 Immunology, Serology and Transplantation

037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

AB Moraxella catarrhalis (formerly known as Branhamella catarrhalis) has emerged as a significant bacterial pathogen of humans over the past two decades. During this period, microbiological and molecular diagnostic techniques have been developed and improved for M.

catarrhalis, allowing the adequate determination and taxonomic positioning of this pathogen. Over the same period, studies have revealed its involvement in respiratory (e.g., sinusitis, otitis media, bronchitis, and pneumonia) and ocular infections in children and in laryngitis, bronchitis, and pneumonia in adults. The development of (molecular) epidemiological tools has enabled the national and international distribution of M. catarrhalis strains to be established, and has allowed the monitoring of nosocomial infections and the dynamics of carriage. Indeed, such monitoring has revealed an increasing number of .beta.-lactamase-positive M. catarrhalis isolates (now well above 90%), underscoring the pathogenic potential of this organism. Although a number of putative M. catarrhalis virulence factors have been identified and described in detail, their relationship to actual bacterial adhesion, invasion, complement resistance, etc. (and ultimately their role in infection and immunity), has been established in a only few cases. In the past 10 years, various animal models for the study of ${\tt M.}$ catarrhalis pathogenicity have been described, although not all of these models are equally suitable for the study of human infection. Techniques involving the molecular manipulation of M. catarrhalis genes and antigens are also advancing our knowledge of the host response to and pathogenesis of this bacterial species in humans, as well as providing insights into possible vaccine candidates. This review aims to outline our current knowledge of M. catarrhalis, an organism that has evolved from an emerging to a well-established human pathogen.

L13 ANSWER 11 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2002-352536 [38] WPIDS

DOC. NO. CPI:

C2002-100176

TITLE:

New Streptococcus protein for the

treatment or prevention of infection or disease

caused by Streptococcus bacteria, such as

meningitis, and for detecting a compound that binds

to the protein.

DERWENT CLASS:

B04 C06 D16

INVENTOR(S):

FRASER, C; GRANDI, G; MARGARIT Y ROS, I; MASIGNANI,

V; TELFORD, J; TETTELIN, H

PATENT ASSIGNEE(S):

(CHIR-N) CHIRON SPA; (GENO-N) INST GENOMIC RES

COUNTRY COUNT:

PATENT INFORMATION:

WEEK LA PG PATENT NO KIND DATE ______

WO 2002034771 A2 20020502 (200238)* EN

98

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TR TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CO CR CU CZ

DE DK DM DZ EC EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP

KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ

NO NZ OM PH PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA

UG US UZ VN YU ZA ZW

APPLICATION DETAILS:

APPLICATION DATE PATENT NO KIND WO 2002034771 A2 WO 2001-GB4789 20011029

PRIORITY APPLN. INFO: GB 2001-5640 20010307; GB 2000-26333 20001027; GB 2000-28727 20001124

AN 2002-352536 [38] WPIDS

AB

WO 200234771 A UPAB: 20020618

NOVELTY - A protein (I) from group B streptococcus (Streptococcus agalactiae) or group A streptococcus (Streptococcus pyogenes), comprising one of 5483 sequences (S1), given in the specification, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) a protein having 50 % or greater sequence
 identity to (I);
- (2) a **protein** comprising a fragment of 7 or more consecutive amino acids from (S1);
 - (3) an antibody which binds (I);
 - (4) a nucleic acid encoding (I);
- (5) a nucleic acid comprising one of 1057 sequences (S2), given in the specification;
- (6) a nucleic acid comprising a fragment of 10 or more consecutive nucleotides from one of 6540 sequences (S3), given in the specification, which includes the sequences of (S2);
- (7) a nucleic acid comprising a sequence complementary to one of (4) (6);
- (8) a nucleic acid comprising a sequence having 50 % or greater sequence identity to one of (4) (7);
- (9) a nucleic acid that can hybridize to (4) (8), under high stringency conditions;
 - (10) a composition comprising (I), or one of (1) (9);
- (11) the use of (10) in the manufacture of a medicament for the treatment of prevention of infection or disease caused by streptococcus bacteria, particularly S. agalactiae and S. pyrogenes;
 - (12) treating a patient comprising administering (10);
 - (13) a hybrid protein of formula (F);
- (14) a kit comprising primers for amplifying a template sequence contained within a Streptococcus nucleic acid sequence, where the kit comprises one primer complementary to the template sequence and a second primer complementary to a complement of the template sequence, and the parts of the primers which have complementarity define the termini of the template sequence to be amplified;
- (15) a kit comprising two single-stranded oligonucleotides which allow amplification of a Streptococcus template nucleic acid contained in a single- or double-stranded nucleic acid (or mixture of it) where:
- (a) one oligonucleotide comprises a primer sequence complementary to the template nucleic acid sequence;
- (b) the second oligonucleotide comprises a primer sequence complementary to the complement of the template nucleic acid sequence;
- (c) either or both oligonucleotides comprise a sequence(s) not complementary to the template nucleic acid sequence; and
- (d) the primer sequences define the termini of the template sequence to be amplified;
- (16) a computer readable medium containing one of 12024 sequences (S4), given in the specification;
 - (17) detecting Streptococcus in a biological sample comprising

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contacting (4) - (9) with the sample under hybridizing conditions;
          (18) determining whether a compound binds to (I), (1), or (2),
     comprising contacting a test compound with the protein and
     determining binding;
          (19) a compound identified by (18);
          (20) a composition comprising (1), (1), or (2) and one of:
          (i) a protein antigen from Helicobacter pylori and/or
     Neisseria meningitidis serogroup B;
          (ii) an outer-membrane vesicle (OMV) preparation from N.
    meningitidis serogroup B;
          (iii) a saccharide antigen from N. meningitidis serogroup A, C,
    W135 and/or Y, or Streptococcus pneumoniae;
          (iv) an antigen from hepatitis A, B, or C virus, and/or
     Bordetella pertussis;
          (v) a diphtheria and/or tetanus antigen;
          (vi) a saccharide antigen from Haemophilus influenzae B;
          (vii) an antigen from N. gonorrhoeae, Chlamydia pneumoniae, C.
     trachomatis, and/or Porphyromonas gingivalis;
          (viii) a polio and/or rabies antigen(s);
          (ix) measles, mumps, and/or rubella antigens;
          (x) an influenza antigen(s);
          (xi) an antigen from Moraxella catarrhalis; and/or
          (xii) an antigen from Staphlococcus aureus; and
          (21) a composition comprising two or more proteins of (1), (1),
     or (2).
         NH2-A-(-X-L-)n-B-COOH
                                  (F)
     X = (I);
         L = an optional linker amino acid sequence;
         A = an optional N-terminal amino acid sequence;
          B = an optional C-terminal amino acid sequence; and
          n = an integer greater than 1.
          ACTIVITY - Antibacterial; antiinflammatory. No suitable
     biological data is given.
          MECHANISM OF ACTION - Gene therapy; vaccine.
          USE - (I), nucleic acids encoding (I), and antibodies that bind
     (I) are used in the manufacture of medicaments for the treatment of
     prevention or infection or disease caused by Streptococcus bacteria,
     particularly S. agalactiae and S. pyrogenes. Nucleic acid encoding
     (I) is used to detect Streptococcus in a biological sample. (I) is
     used to determine whether a compound binds to (I). A composition
     comprising (I) or a nucleic acid encoding (I), may be used as a
     vaccine or diagnostic composition (all claimed). The disease caused
     by Streptococcus that is prevented or treated may be meningitis.
     Nucleic acid encoding (I) may be used to recombinantly produce (I).
     Antibodies to (I) are used for affinity chromatography,
     immunoassays, and distinguishing/identifying Streptococcus proteins.
     Dwg.0/319
                      WPIDS (C) 2002 THOMSON DERWENT
L13 ANSWER 12 OF 37
                      2001-244783 [25]
                                         WPIDS
ACCESSION NUMBER:
                      N2001-174285
DOC. NO. NON-CPI:
                      C2001-073454
DOC. NO. CPI:
                      Novel BASB129-BASB131 polypeptides
TITLE:
                      isolated from Moraxella catarrhalis bacterium
                      useful as a diagnostic reagent for M.catarrhalis
                      infections and for producing vaccines against
                      otitis media and pneumonia.
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Searcher: Shears 308-4994

B04 D16 S03

DERWENT CLASS:

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

95

PATENT INFORMATION:

PATENT NO	KIND DATE	WEEK	LA	PG

WO 2001019862 A2 20010322 (200125)* EN 80

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

AU 2001013839 A 20010417 (200140)

EP 1214339 A2 20020619 (200240) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND .	API	PLICATION	DATE
WO 2001019862 AU 2001013839 EP 1214339		AU EP	2000-EP9034 2001-13839 2000-975853 2000-EP9034	20000914 20000914 20000914 20000914

FILING DETAILS:

	KIND	PATENT NO
AU 20010138	39 A Based on	WO 200119862 WO 200119862

PRIORITY APPLN. INFO: GB 1999-22829 19990925; GB 1999-21693 19990914; GB 1999-21694 19990914

AN 2001-244783 [25] WPIDS

AB WO 200119862 A UPAB: 20010508

NOVELTY - Isolated Moraxella catarrhalis BASB129-BASB131 polypeptides (I) comprising a fully defined sequence of 344 (S2), 678 (S4), 469 (S6) amino acids, respectively as given in the specification, or an isolated polypeptide (Ia) which has 85% identity to (S2), (S4) or (S6), is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an immunogenic fragment (II), of (I) which has the same immunogenic activity as (I);
- (2) an isolated polynucleotide (III), or its complementary nucleotide sequence comprising a nucleotide sequence:
- (i) encoding a polypeptide that has 85% identity over the entire length of (S2), (S4), (S6);
- (ii) that has 85% identity over the entire length of the nucleotide sequence encoding region which encodes (S2), (S4), (S6);
- (iii) which has 85% identity over the entire length of a fully defined nucleotide sequence of 1035 (S1), 2037 (S3), 1410 (S5), base pairs as given in the specification;

- (iv) comprising a nucleotide sequence encoding (I) obtainable by screening an appropriate library under stringent hybridization conditions with a labeled probe having the sequence of (S1), (S3), (S5);
 - (v) encoding (S2), (S4) or (S6); or
 - (vi) an isolated polynucleotide comprising (S1), (S3) or (S5);
- (3) an expression vector (IV), or a recombinant live microorganism comprising (III);
- (4) a host cell (V) comprising (IV), or a subcellular fraction or membrane of the host cell expressing (I);
 - (5) preparation of (I) or (II);
- (6) expressing (III) involves transforming (V) with (IV) which contains any one of the polynucleotides (III) given above and culturing (V) under suitable conditions to express (III);
 - (7) a vaccine composition which comprises (I) or (II);
 - (8) a vaccine composition which comprises (III);
 - (9) an antibody (Ab) immunospecific for (I) or (II);

and

(10) a therapeutic composition comprising an **antibody** directed against (I) useful in treating humans with M.catarrhalis disease.

ACTIVITY - Antiinflammatory; auditory.

MECHANISM OF ACTION - Gene therapy; vaccine; initial physical attraction between a pathogen and a mammalian extracellular matrix protein inhibitor.

The biological activity of (I) was tested in mice. Groups of mice were immunized with BASB129, BASB130 and BASB131 vaccine. After the booster, the mice were challenged by bacterial suspension into the nostril under anesthesia. Mice were killed between 30 minutes and 24 hours after challenge and the lungs were removed and homogenized. The log10 weighted mean number of colony forming unit (CFU)/lung was determined by counting the colonies grown on agar plates after plating of dilutions of the homogenate. The arithmetic mean of the log10 weighted mean number of CFU/lung and the standard deviations were calculated for each group. Results were analyzed statistically. Results showed that BASB129, BASB130 and BASB131 vaccine induced significant lung clearance as compared to the control group.

USE - The composition comprising (I), (II) or (III) is useful for preparation of a medicament used for generating an immune response in an animal. (I) is also useful as diagnostic reagent for M.catarrhalis which involves identifying (I), an antibody against (I) present within the biological sample from an animal suspected of having such an infection (claimed). Fragments of (I) are useful for producing corresponding full length polypeptides by peptide synthesis. The polynucleotides may be used as hybridization probes for RNA, cDNA and genomic DNA to isolate full-length cDNAs and genomic clones encoding BASB129-BASB131 and to isolate cDNA and genomic clones of other genes that have high sequence identity to BASB129-BASB131 gene. The polynucleotide sequences can also be used in the discovery and development of antibacterial compounds. The encoded protein can be used as target for the screening of antibacterial drugs. Additionally, the polynucleotide sequences encoding the amino terminal regions of the encoded protein or Shine-Dalgarno or other translation facilitating sequences of the respective mRNA can be used to construct antisense sequences to control the expression of the coding sequence of interest. The

polynucleotides are also useful as diagnostic reagents in which the mutations in the polynucleotide sequence may be detected and used to diagnose and/or prognose the infection or its stage or course. The polynucleotides may be used as components of arrays which have diagnostic and prognostic uses. Antibodies against (I) are useful for treating bacterial infections and to isolate or identify clones expressing (I) or (II), to purify the polypeptides by affinity chromatography. The polynucleotides and polypeptides are used as research reagents and materials for discovery of treatments of and diagnostics for human diseases. The polynucleotides derived from (S1), (S3) or (S5) are used as PCR (polymerase chain reaction) primers. The polynucleotides are also useful in the diagnosis of the stage of infection and type of infection the pathogen has attained. The polypeptides and polynucleotides are used to block the initial physical interaction between a gram negative and/or gram positive bacteria to mammalian, host thus preventing the sequelae of infection. The polynucleotides encoding certain non-variable regions of bacterial cell surface protein are used in polynucleotide constructs which are useful for genetic immunization experiments in animal models of infection with M.catarrhalis to identify protein groups able to provoke a prophylactic or therapeutic immune response. The vaccine comprising (I), (II) or (III) is useful for treating Moraxella catarrhalis infections such as sinusitis, nosocomial infections, otitis media and pneumonia. Dwq.0/0

L13 ANSWER 13 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2001-182955 [18] WPIDS

DOC. NO. NON-CPI:

N2001-130566

DOC. NO. CPI:

C2001-054636

TITLE:

New BASB126 polypeptides of Moraxella

catarrhalis useful for diagnostic, prophylactic and therapeutic purposes against microbial diseases,

preferably bacterial infections.

DERWENT CLASS:

B04 D16 S03

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

95

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001009329 A1 20010208 (200118)* EN 86

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

AU 2000068316 A 20010219 (200129) EP 1204750 A1 20020515 (200239) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL RO SI

APPLICATION DETAILS:

PATENT NO KIND	APPLICATION	DATE
WO 2001009329 A1 AU 2000068316 A EP 1204750 A1	WO 2000-EP7280 AU 2000-68316 EP 2000-956332 WO 2000-EP7280	20000727 20000727 20000727 20000727

FILING DETAILS:

PATENT NO K	IND	PATENT NO
AU 2000068316	A Based on	WO 200109329
EP 1204750	Al Based on	WO 200109329

PRIORITY APPLN. INFO: GB 1999-18038 19990730

AN 2001-182955 [18] WPIDS

AB WO 200109329 A UPAB: 20010402

NOVELTY - An isolated BASB126 polypeptide (I) of Moraxella catarrhalis, comprises a sequence having at least 85% identity (over the entire length) to one of the two 192 amino acids sequences given in the specification.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an immunogenic fragment (II) of (I), where (II) has the same immunogenicity of (I);
 - (2) an isolated polynucleotide (III) encoding (I) (II);
- (3) an expression vector (IV) or a recombinant live microorganism, comprising (III);
- (4) a host cell (V) comprising (IV), or a subcellular fraction or membrane of (V) expressing (I);
- (5) producing (I) comprising culturing (V) and recovering the polypeptide from the culture medium;
- (6) expressing (III) comprising transforming (V) with (IV) and culturing under conditions sufficient for its expression;
 - (7) a vaccine (VI) comprising (I), (II) or (III);
 - (8) an antibody (VII) immunospecific for (I) or (II);
- (9) diagnosing Moraxella catarrhalis infection comprising identifying (I) or (VII) in a biological sample from an animal suspected of having such an infection; and
- (10) a therapeutic composition (VIII) for treating Moraxella catarrhalis infection comprising at least one (VII).

ACTIVITY - Antibacterial; antimicrobial; auditory; antiinflammatory.

MECHANISM OF ACTION - Vaccine.

Experimental protocols are described but no results are given. USE - (VI) is useful for preparing a medicament for use in generating immune response in an animal (claimed). (VIII) is useful for treating humans with Moraxella catarrhalis disease (claimed).

(I) and (III) are useful in the prevention, treatment and diagnosis of microbial diseases, preferably bacterial infections such as otitis media, pneumonia, sinusitis, nosocomial infections, and invasive diseases. (I) and (III) are useful as immunogens to produce antibodies, and to asses the binding of small molecule substrate and ligands in, for e.g., cells, cell-free preparations, chemical libraries and natural product mixtures. (I), (III) and (VII) are useful to configured screening methods for detecting the effect of added compounds and production of mRNA and/or polypeptides in the cells.

(III) is useful as a hybridization probe for RNA, cDNA and genomic DNA to isolate full-length cDNAs and genomic clones encoding BASB126 and to isolate cDNA and genomic clones of other genes that have a high identity particularly high sequence identity, to the BASB126 gene. (II) has utility in diagnosis of the stage and type of infection, and also for therapeutic or prophylactic purposes, in particular genetic immunization. (II) is useful as a component of polynucleotide arrays, preferably high density arrays or grid. Dwg.0/4

L13 ANSWER 14 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2001-168707 [17] WPIDS

DOC. NO. NON-CPI:

N2001-121639

DOC. NO. CPI:

C2001-050432

TITLE:

New BASB125 polypeptide isolated from

Moraxella catarrhalis for treating, preventing and diagnosing diseases associated with M. catarrhalis infection in mammals, e.g. otitis media in humans.

DERWENT CLASS:

B04 D16 S03

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

95

PATENT INFORMATION:

PATENT	NO	KIND	DATE	WEEK	LA	PG
				-		

WO 2001009331 A2 20010208 (200117) * EN 73

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000064393 A 20010219 (200129)

EP 1212424 A2 20020612 (200239) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO KI	IND	APE	PLICATION	DATE
WO 2001009331 AU 2000064393 EP 1212424		AU EP	2000-EP7291 2000-64393 2000-951466 2000-EP7291	20000727 20000727 20000727 20000727

FILING DETAILS:

PATENT NO ·K	IND	PATENT NO
AU 2000064393 EP 1212424		WO 200109331 WO 200109331

PRIORITY APPLN. INFO: GB 1999-18041

19990730

AN 2001-168707 [17] WPIDS

AB WO 200109331 A UPAB: 20010328

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Searcher :

Shears 308-4994

NOVELTY - An isolated **polypeptide** having at least 85 % identity to a sequence (I) of 134 amino acids for a Moraxella catarrhalis BASB125 **polypeptide**, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

(1) an isolated polypeptide of sequence (I);

(2) immunogenic fragments of the polypeptide having the same immunogenic activity as sequence (I);

(3) an isolated polynucleotide:

(i) having 85 % identity to a polynucleotide encoding the polypeptide, especially 85 % identity to sequence (II) of 405 base pairs (bp) encoding sequence (I);

(ii) complementary to a polynucleotide of (i);

(iii) encoding the new polypeptide; and

- (iv) encoding sequence (I) and obtained by screening a library under stringent conditions using sequence (II) or a fragment as a probe;
- (4) vectors or recombinant live microorganisms comprising the polynucleotide;
- (5) host cells comprising the vector and subcellular fragments/membranes of the host cells expressing the polypeptide;
- (6) producing the new polypeptide comprising culturing the host cell of (5) to produce the polypeptide and recovering the polypeptide from the culture medium;
- (7) expressing (3) comprising transforming a host cell with an expression vector of (4) and culturing the host cell to express the polynucleotide;
- (8) vaccine compositions comprising the new **polypeptide** or (3):

(9) antibodies specific for the new polypeptide, or immunological fragments of (2);

(10) diagnosing a M. catarrhalis infection comprising identifying the new polypeptide or an antibody immunospecific for the polypeptide, present within a biological sample from an animal suspected of having the infection;

(11) preparing a medicament for generating an immune response in an animal using a composition comprising the new polypeptide or (3); and

(12) a therapeutic composition for treating humans with M.catarrhalis disease comprising an antibody against the new polypeptide.

ACTIVITY - Antibacterial. A sequence (II) of 405 base pairs (bp) was isolated from M. catarrhalis strain American Type Culture Collection (ATCC) 43617 by standard molecular biological techniques a sequence (I) of 134 amino acids deduced. Mice were immunized with a BASB125 vaccine or a killed whole cell (kwc) M. catarrhalis preparation, or were sham immunized. After a booster, mice were challenged by instillation of bacterial suspension into the nostril under anaesthesia. Mice were killed 30 minutes-24 hours after challenge and lungs removed aseptically and homogenized. Homogenates were diluted and plated onto agar plates, and log10 weighted mean number of colony forming units/lung determined by counting. BASB125 vaccine and kwc preparations induced significant lung clearance of M. catarrhalis versus controls. No experimental data is given.

MECHANISM OF ACTION - Vaccine; gene therapy.

USE - The polypeptide, immunogenic fragments of the polypeptide, fusion proteins of the

polypeptide, or polynucleotides encoding the polypeptide are used in vaccine compositions (claimed), optionally with another M. catarrhalis antigen (claimed). They can also be included in medicaments for use in generating an immune response in an animal (claimed). The vaccines and medicaments are useful in preventing and/or treating microbial diseases, especially diseases associated with M. catarrhalis infection in mammals (especially humans). The polypeptides/polynucleotides may be used to produce antibodies, which can be used in compositions useful therapeutically to treat humans with M. catarrhalis diseases (claimed). M. catarrhalis is a Gram-negative bacteria frequently isolated from the human upper respiratory tract and responsible for several pathologies in humans e.g. otitis media in children, pneumonia, sinusitis etc. The polypeptides, polynucleotides and antibodies are also useful diagnostically e.g. in the detection of the polypeptides/ antibodies in a biological sample from an animal to diagnose M. catarrhalis infection (claimed). The diagnostic assays are useful e.g. to detect diseases, determine the stage and type of infection, determine the effect of drugs etc. The polypeptides and polynucleotides can also be used to detect antagonists and agonists useful e.g. in preventing, inhibiting and/or treating disease. The polynucleotides are also useful in producing hybridization probes to isolate sequences encoding BASB125 and similar sequences. Dwg.0/0

L13 ANSWER 15 OF 37 WPIDS (C) 2002 THOMSON DERWENT WPIDS

2001-159876 [16] ACCESSION NUMBER:

N2001-116486 DOC. NO. NON-CPI: C2001-047628 DOC. NO. CPI:

New BASB117 polypeptides from Moraxella TITLE:

catarrhalis strain MC2931 (ATCC 43617), useful as therapeutic agents or vaccines against bacterial (especially M. catarrhalis) infections, e.g. otitis

media or pneumonia.

B04 D16 S03 DERWENT CLASS: INVENTOR(S): THONNARD, J

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS PATENT ASSIGNEE(S):

COUNTRY COUNT: 95

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001009339 A2 20010208 (200116) * EN

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

AU 2000065688 A 20010219 (200129) EP 1206547 A2 20020522 (200241) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	API	PLICATION	DATE
WO 2001009339 AU 2000065688 EP 1206547	•	AU EP	2000-EP7422 2000-65688 2000-953131 2000-EP7422	20000731 20000731 20000731 20000731

FILING DETAILS:

PAT	TENT NO	KIND			PAT	TENT NO
AU	200006568	 8 A	Based	on	WO	200109339
EΡ	1206547	A2	Based	on	WO	200109339

PRIORITY APPLN. INFO: GB 1999-18206 19990803

AN 2001-159876 [16] WPIDS

AB WO 200109339 A UPAB: 20010323

NOVELTY - Moraxella catarrhalis strain MC2931 (ATCC 43617) BASB117 polypeptides, both of 216 amino acids (I and II) as defined in the specification, are new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated **polypeptide** (P1) comprising an amino acid sequence which has at least 85%, preferably 100%, identity to (I) or (II) over their entire length;
- (2) an immunogenic fragment (P2) of the polypeptide, in which the immunogenic activity of the fragment is substantially the same as (I) or (II);
 - (3) an isolated polynucleotide (N1) selected from:
 - (a) a nucleotide sequence encoding (I), (II), P1 or P2;
- (b) an isolated polynucleotide comprising a nucleotide sequence encoding a **polypeptide** that has at least 85%, preferably 95%, identity to (I) or (II) over its entire length, or a nucleotide sequence complementary to the isolated polynucleotide;
- (c) an isolated polynucleotide comprising a nucleotide sequence that has at least 85%, preferably 95%, identity to a nucleotide sequence encoding (I) or (II) over the entire coding region, or a nucleotide sequence complementary to the isolated polynucleotide;
- (d) an isolated polynucleotide comprising the 648 (III) or 651 basepair (bp) sequence (IV) fully defined in the specification;
- (e) an isolated polynucleotide comprising a nucleotide sequence which has at least 85%, preferably 95%, identity to (I) or (II) over its entire length, or a nucleotide sequence complementary to the isolated polynucleotide;
- (f) a nucleotide sequence encoding (I) or (II) obtainable by screening an appropriate library, under stringent conditions, with a labeled probe having the sequence of (III), (IV) or its fragments;
- (4) an expression vector or a recombinant live microorganism comprising N1;
- (5) a host cell comprising the expression vector of (4), or a subcellular fraction or membrane of the host cell expressing P1;
- (6) a process for producing (I), (II), P1 or P2 by culturing the host cell of (5);
- (7) a process for expressing N1 comprising transforming a host cell with the expression vector of (4) and culturing the host cell;
 - (8) a vaccine compositions comprising (I), (II), P1 or P2 or

N1;

(9) an antibody immunospecific for (I), (II), Pl or P2;

(10) a method for diagnosing a Moraxella catarrhalis infection comprising identifying (I), (II), P1 or P2 or the **antibody** of (9) present within a biological sample from an animal suspected of having such an infection; and

(11) a therapeutic composition for treating humans with Moraxella catarrhalis disease, comprising at least one antibody against (I), (II), P1 or P2.

ACTIVITY - Antibacterial; ophthalmological; antiinflammatory.

MECHANISM OF ACTION - Vaccine; gene therapy.

Groups of mice were immunized with the polypeptide

(BASB117) or with a killed whole cells (kwc) preparation of Moraxella catarrhalis or sham immunized.

After booster, the mice were challenged by instillation of bacterial suspension into the nostril under anaesthesia. Mice were killed between 30 minutes and 24 hours after challenge and the lungs were removed aseptically and homogenized individually. The log10 weighted mean number of colony forming units (CFU)/lung was determined by counting the colonies grown on agar plates after plating of dilutions of the homogenate. The arithmetic mean of the log10 weighted mean number of CFU/lung and the standard deviations were calculated for each group.

No results are given.

USE - The composition comprising an immunologic amount of the polypeptide or polynucleotide is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection (all claimed). The polypeptides may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptides or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the polypeptides or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging of the disease, determining the response of an infectious organism to drugs. Dwg.0/2

L13 ANSWER 16 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2001-159875 [16] WPIDS

DOC. NO. NON-CPI:

N2001-116485 C2001-047627

DOC. NO. CPI: TITLE:

New BASB116 polypeptides from Moraxella

catarrhalis strain MC2931 (ATCC 43617), useful as therapeutic agents or vaccines against bacterial (especially M. catarrhalis) infections, e.g. otitis

media or pneumonia.

DERWENT CLASS:

B04 D16 S03

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

95

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001009338 A1 20010208 (200116) * EN 79

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000062788 A 20010219 (200129)

EP 1206545 A1 20020522 (200241) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	API	PLICATION	DATE
WO 2001009338 AU 2000062788 EP 1206545		AU EP	2000-EP7421 2000-62788 2000-949429 2000-EP7421	20000731 20000731 20000731 20000731

FILING DETAILS:

PATENT NO K	 PATENT NO
AU 2000062788 EP 1206545	WO 200109338 WO 200109338

PRIORITY APPLN. INFO: GB 1999-18279 19990803

AN 2001-159875 [16] WPIDS

AB WO 200109338 A UPAB: 20010323

NOVELTY - Two Moraxella catarrhalis strain MC2931 (ATCC 43617) BASB116 polypeptides, both of 98 amino acids (I and II) as defined in the specification, are new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated **polypeptide** (P1) comprising an amino acid sequence which has at least 85%, preferably 100%, identity to (I) or (II) over their entire length;
- (2) an immunogenic fragment (P2) of the polypeptide, in which the immunogenic activity of the fragment is substantially the same as (I) or (II);
 - (3) an isolated polynucleotide (N1) selected from:
 - (a) a nucleotide sequence encoding (I), (II), P1 or P2;
- (b) an isolated polynucleotide comprising a nucleotide sequence encoding a polypeptide that has at least 85% identity to
 (I) or (II) over its entire length, or a nucleotide sequence complementary to the isolated polynucleotide;
- (c) an isolated polynucleotide comprising a nucleotide sequence that has at least 85%, preferably 95%, identity to a nucleotide

- sequence encoding (I) or (II) over the entire coding region, or a nucleotide sequence complementary to the isolated polynucleotide;
- (d) an isolated polynucleotide comprising the 297 (III) or 294 (IV) basepair (bp) sequence fully defined in the specification;
- (e) an isolated polynucleotide comprising a nucleotide sequence which has at least 85%, preferably 95%, identity to (I) or (II) over its entire length, or a nucleotide sequence complementary to the isolated polynucleotide;
- (f) a nucleotide sequence encoding (I) or (II) obtainable by screening an appropriate library, under stringent conditions, with a labeled probe having the sequence of (III), (IV) or its fragments;
- (4) an expression vector or a recombinant live microorganism comprising N1;
- (5) a host cell comprising the expression vector of (4), or a subcellular fraction or membrane of the host cell expressing P1;
- (6) a process for producing (I), (II), P1 or P2 by culturing
 the host cell of (5);
- (7) a process for expressing N1 comprising transforming a host cell with the expression vector of (4) and culturing the host cell;
- (8) a vaccine compositions comprising (I), (II), P1 or P2 or N1;
- (9) an antibody immunospecific for (I), (II), Pl or P2;
- (10) a method for diagnosing a Moraxella catarrhalis infection comprising identifying (I), (II), P1 or P2 or the **antibody** of (9) present within a biological sample from an animal suspected of having such an infection; and
- (11) a therapeutic composition for treating humans with Moraxella catarrhalis disease, comprising at least one antibody against (I), (II), Pl or P2.

ACTIVITY - Antibacterial; ophthalmological; antiinflammatory. MECHANISM OF ACTION - Vaccine; gene therapy.

Groups of mice were immunized with the **polypeptide** (BASB116) or with a killed whole cells (kwc) preparation of Moraxella catarrhalis or sham immunized.

After booster, the mice were challenged by instillation of bacterial suspension into the nostril under anaesthesia. Mice were killed between 30 minutes and 24 hours after challenge and the lungs were removed aseptically and homogenized individually. The log10 weighted mean number of colony forming units (CFU)/lung was determined by counting the colonies grown on agar plates after plating of dilutions of the homogenate. The arithmetic mean of the log10 weighted mean number of CFU/lung and the standard deviations were calculated for each group.

No results are given.

USE - The composition comprising an immunologic amount of the polypeptide or polynucleotide is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection (all claimed). The polypeptides may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory

tract, or inflammation of the middle ear. The polypeptides or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the polypeptides or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging of the disease, determining the response of an infectious organism to drugs.

Dwg.0/2

L13 ANSWER 17 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2001-159874 [16] WPIDS

DOC. NO. NON-CPI: N2001-116484 DOC. NO. CPI: C2001-047626

TITLE: New BASB122 and BASB124 polypeptides and

polynucleotides from Moraxella catarrhalis strain

ATCC 43617, useful as therapeutic agents or

vaccines against bacterial infections, e.g. otitis

media or pneumonia.

DERWENT CLASS:

B04 D16 S03 THONNARD, J

INVENTOR(S):
PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT: 95

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001009337 A2 20010208 (200116)* EN 75

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

AU 2000065683 A 20010219 (200129)

EP 1204749 A2 20020515 (200239) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	API	PLICATION	DATE
WO 2001009337 AU 2000065683 EP 1204749		AU EP	2000-EP7365 2000-65683 2000-953120 2000-EP7365	20000731 20000731 20000731 20000731

FILING DETAILS:

PAT	ENT	NO	KIND			 ENT	
			3 A	Based	on	 2001	.09337

PRIORITY APPLN. INFO: GB 1999-18036 19990730; GB 1999-18034

19990730

AN 2001-159874 [16] WPIDS

AB

WO 200109337 A UPAB: 20010323

NOVELTY - New isolated **polypeptides**, comprising either of two 111 amino acid (I) or two 328 amino acid (II) sequences from Moraxella catarrhalis, all fully defined in the specification, or an at least 85 % identical sequence over their entire length, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

(1) an isolated polynucleotide encoding the novel polypeptide, comprising:

(a) a sequence encoding the novel polypeptide;

- (b) a sequence having at least 85 % identity to (a) over its entire length;
- (c) a 336 (III) or 987 (IV) base pair sequence, both fully defined in the specification;
- (d) a sequence at least 85 % identical to (III) or (IV) over their entire length;

(e) the complements of (a)-(d); or

- (f) a sequence encoding (I) or (II) obtained by screening a library, under stringent conditions, with a labeled probe having (III), (IV), or fragments of them;
- (2) a statement vector or a recombinant live microorganism, comprising the polynucleotide of (1);
- (3) a host cell comprising the vector of (2), or a subcellular fraction or membrane of the host cell expressing the novel polypeptide;
- (4) a process for producing the novel **polypeptide**, comprising culturing the host cell of (3) under expression conditions, and recovering the **polypeptide**;
- (5) a process for expressing the polynucleotide of (1), comprising transforming a host cell with the vector of (2), and culturing the cell for expression of the polynucleotide;
- (6) a vaccine composition comprising the novel polypeptide or the polynucleotide of (1), and a carrier;
- (7) an antibody immunospecific for the novel polypeptide or its immunological fragment;
- (8) a method for diagnosing a M. catarrhalis infection, comprising identifying the novel **polypeptide** or the **antibody** of (7) present within a biological sample; and
- (9) a therapeutic composition comprising at least one antibody against the novel polypeptide.

ACTIVITY - Antibacterial; antiinflammatory; auditory. MECHANISM OF ACTION - Vaccine; gene therapy.

No biological data is given.

USE - The composition comprising an immunologic amount of the polypeptide or polynucleotide is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection. (All claimed). The polypeptides may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptides

or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the polypeptides or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging of the disease, determining the response of an infectious organism to drugs. Dwg.0/0

L13 ANSWER 18 OF 37

WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2001-159873 [16] WPIDS

DOC. NO. NON-CPI:

N2001-116483

DOC. NO. CPI:

C2001-047625

TITLE:

New BASB119 polypeptides and

polynucleotides from Moraxella catarrhalis strain

ATCC 43617, useful as therapeutic agents or

vaccines against bacterial infections, e.g. otitis

media or pneumonia.

DERWENT CLASS:

B04 D16 S03

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

95

PATENT INFORMATION:

PATENT	NO	KIND	DATE	WEEK	LA	PG

WO 2001009336 A1 20010208 (200116) * EN 82

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

AU 2000069887 A 20010219 (200129)

EP 1206549 A1 20020522 (200241) EN

> R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	API	PLICATION	DATE
WO 2001009336 AU 2000069887 EP 1206549		AU EP	2000-EP7363 2000-69887 2000-958324 2000-EP7363	20000731 20000731 20000731 20000731

FILING DETAILS:

PATENT NO	KIND	PATENT NO
AU 200006988	7 A Based on	WO 200109336
EP 1206549	Al Based on	WO 200109336

PRIORITY APPLN. INFO: GB 1999-18302

19990803

AN 2001-159873 [16] WPIDS

AB WO 200109336 A UPAB: 20010323

> 308-4994 Searcher : Shears

NOVELTY - New isolated **polypeptides**, comprising either of two 171 residue amino acid sequences (I and II) from Moraxella catarrhalis, both fully defined in the specification, or a sequence at least 85 % identical to (I) or (II), over their entire length, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated polynucleotide encoding the novel polypeptide, comprising:
 - (a) a sequence encoding (I) or (II);
- (b) a sequence having at least 85 % identity to the sequence encoding (I) or (II) over the entire coding region;
- (c) a 516 (III) or 513 (IV) base pair sequence, fully defined in the specification;
- (d) a sequence having at least 85 % identity to (III) or (IV) over their entire length;
 - (e) the complements of (a)-(d); or
- (f) a sequence encoding (I) or (II) obtained by screening a library, under stringent conditions, with a labeled probe having (III), (IV), or fragments of (III) or (IV);
- (2) an statement vector or a recombinant live microorganism comprising the polynucleotide of (1);
- (3) a host cell comprising the vector of (2), or a subcellular fraction or membrane of the host cell expressing the novel polypeptide;
- (4) a process for producing the novel **polypeptide**, comprising culturing the cell of (3) under expression conditions, and recovering the **polypeptide**;
- (5) a process for expressing the polynucleotide of (1), comprising transforming a host cell with the vector of (2), and culturing the host cell for expression of the polynucleotide;
- (6) vaccine compositions comprising the novel polypeptide or the polynucleotide of (1), and a carrier;
- (7) an antibody immunospecific for the novel
- polypeptide or its immunological fragment;
- (8) a method for diagnosing a M. catarrhalis infection, comprising identifying the novel **polypeptide** or the **antibody** present within a biological sample; and
- (9) a therapeutic composition comprising at least one antibody against the novel polypeptide.

ACTIVITY - Antibacterial; antiinflammatory; auditory. MECHANISM OF ACTION - Vaccine; gene therapy.

Groups of mice were immunized either with the polypeptide (BASB119) adsorbed onto AlPO4 (10 micro g BASB119 onto 100 micro g of AlPO4), with a killed whole cell (kwc) preparation of M. catarrhalis strain ATCC 43617 adsorbed onto AlPO4, or with 100 micro g AlPO4 without antigen. The mice were challenged with 5 multiply 105 colony forming units (CFU) of live M. catarrhalis strain ATCC 43617 bacteria. The log10 weighted mean number of CFU/lung and the standard deviation 4 hours after challenge was calculated for each group. Sham immunized mice had 5.41 (+/-0.2) log10 CFU/lungs 4 hours after challenge. The kwc preparation induced significant lung clearance as compared to the control group (1.58 log difference). BASB119 vaccine induced a 1.34 log difference in lung clearance, which was significantly different from the control.

USE - The composition comprising the novel **polypeptide** or polynucleotide is useful for preparing a medicament for

generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection. (All claimed). The polypeptides may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptides or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the polypeptides or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging of the disease, determining the response of an infectious organism to drugs. Dwg.0/3

L13 ANSWER 19 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2001-159872 [16] WPIDS

DOC. NO. NON-CPI: N2001-116482 DOC. NO. CPI: C2001-047624

TITLE: New BASB120 polypeptides and

polynucleotides from Moraxella catarrhalis strain American Type Culture Collection 43617, for use as therapeutic agents or vaccines against bacterial

infections, e.g. otitis media or pneumonia.

DERWENT CLASS: B04 D16 S03 INVENTOR(S): THONNARD, J

PATENT ASSIGNEE(S): (SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT: 95

PATENT INFORMATION:

PATENT	NO	KIND	DATE	WEEK	LA	PG

WO 2001009335 A2 20010208 (200116) * EN 75

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ

PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000064397 A 20010219 (200129)

EP 1206546 A2 20020522 (200241) EN

R: AL AT BE CH CY DE DK ES FI FK GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO KI	IND	APPLICATION	DATE
WO 2001009335 AU 2000064397		WO 2000-EP7361 AU 2000-64397	20000731 20000731
EP 1206546	A2	EP 2000-951472	20000731

WO 2000-EP7361 20000731

FILING DETAILS:

PRIORITY APPLN. INFO: GB 1999-18281 19990803

AN 2001-159872 [16] WPIDS

AB WO 200109335 A UPAB: 20010323

NOVELTY - An isolated polypeptide (PP) comprising:

- (a) a sequence of 250 amino acids (I) from Moraxella catarrhalis, given in the specification; or
- (b) an amino acid sequence, which has at least 85% identity to(I), over the entire length of (I), is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an immunogenic fragment of the polypeptide, inwhich the immunogenic activity of the fragment is the same as (I);
- (2) isolated polynucleotides, which encode the polypeptides, comprising:

(i) a nucleotide sequence encoding (PP);

- (ii) a nucleotide sequence having 85% identity to the nucleotide sequence encoding (I) over the entire coding region;
- (iii) a 753 base pair (bp) DNA sequence (II), given in the specification;
- (iv) a nucleotide sequence having 85% identity to (II) over the entire length of (II);

(v) the complements of (i)-(iv); or

- (vi) a nucleotide sequence encoding (I) obtainable by screening an appropriate library, under stringent conditions, with a labeled probe having (II) or its fragments;
- (3) an expression vector or a recombinant live microorganism comprising (2);
- (4) a host cell comprising the expression vector, or a subcellular fraction or membrane of the host cell expressing (PP);
- (5) producing (PP) comprising culturing (4) to produce (PP) and recovering (PP) from the culture medium;
- (6) expressing (2) comprising transforming a host cell with the expression vector and culturing the host cell for expression of any of the polynucleotides;
- (7) vaccine compositions comprising (PP) or (2), and a pharmaceutical carrier;
- (8) an antibody immunospecific for (PP) or immunological fragment of (1);
- (9) diagnosing a M. catarrhalis infection comprising identifying (PP) or the **antibody** of (8) present within a biological sample from an animal suspected of having such an infection;
- (10) using the compositions of (7) for preparing a medicament for use in generating an immune response in an animal; and
- (11) a therapeutic composition comprising the ${\bf antibody}$ of (8).

ACTIVITY - Antibacterial; antiinflammatory; pulmonary.
MECHANISM OF ACTION - Vaccine; gene therapy. Clinical test
details are described but no results are given.

USE - A composition comprising an immunologic amount of (PP) or a polynucleotide encoding it, is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection (all claimed). The polypeptides may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptides or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the polypeptides or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging diseases, and determining the response of an infectious organism to drugs. Dwg.0/2

L13 ANSWER 20 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2001-159871 [16] WPIDS

DOC. NO. NON-CPI: N2001-116481 DOC. NO. CPI: C2001-047623

TITLE: New BASB118 polypeptides and

polynucleotides from Moraxella catarrhalis strain American Type Culture Collection 43617, for use as therapeutic agents or vaccines against bacterial

infections, e.g. otitis media or pneumonia.

DERWENT CLASS: B04 D16 S03

INVENTOR(S): THONNARD, J

PATENT ASSIGNEE(S): (SMIK) SMITHKLINE BEECHAM BIOLOGICALS COUNTRY COUNT: 95

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001009334 A1 20010208 (200116) * EN 77

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW
W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE

DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG
KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ
PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN
YU ZA ZW

AU 2000068330 A 20010219 (200129) EP 1206548 A1 20020522 (200241) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO KIND APPLICATION DATE

WO 2001009334 A1 WO 2000-EP7360 20000731

AU 2000068330 A EP 1206548 A1 AU 2000-68330 20000731 EP 2000-956353 20000731 WO 2000-EP7360 20000731

FILING DETAILS:

PATENT NO PATENT NO KIND WO 200109334 AU 2000068330 A Based on WO 200109334 Al Based on EP 1206548

19990803 PRIORITY APPLN. INFO: GB 1999-18208

2001-159871 [16] WPIDS AB

WO 200109334 A UPAB: 20010323

NOVELTY - An isolated polypeptide comprising:

- (a) a sequence of 386 amino acids (I) from Moraxella catarrhalis, given in the specification; or
- (b) an amino acid sequence, which has 85% identity to (I), over the entire length of (I), is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an immunogenic fragment of the new polypeptide, in which the immunogenic activity of the fragment is the same as
- (2) isolated polynucleotides, which encode the new polypeptide, comprising:

(i) a nucleotide sequence encoding (a) or (b);

- (ii) a nucleotide sequence that has 85% identity to the nucleotide sequence encoding (I) over the entire coding region;
- (iii) a 1161 base pair (bp) DNA sequence (II), given in the specification;
- (iv) a nucleotide sequence that has 85% identity to (II) over the entire length of (II);

(v) the complements of (i)-(iv); or

- (vi) a nucleotide sequence encoding (I) obtainable by screening an appropriate library, under stringent conditions, with a labeled probe having (II) or its fragments;
- (3) an expression vector or a recombinant live microorganism comprising an isolated polynucleotide of (2);
- (4) a host cell comprising the expression vector of (3), or a subcellular fraction or membrane of the host cell expressing the new polypeptide;
- (5) producing the new polypeptide comprising culturing (4) to produce the new polypeptide and recovering it from the culture medium;
- (6) expressing a polynucleotide of (2) comprising transforming a host cell with the expression vector of (3) and culturing the host cell for expression of any of the polynucleotides;
- (7) vaccine compositions comprising the new polypeptide or polynucleotide of (2), and a pharmaceutical carrier;

(8) an antibody immunospecific for the new polypeptide or immunological fragment;

(9) diagnosing a M. catarrhalis infection comprising

- identifying the new polypeptide or the antibody of (8) present within a biological sample from an animal suspected of having such an infection; and
- (10) a therapeutic composition comprising an antibody of (8).

ACTIVITY - Antibacterial; antiinflammatory; pulmonary. MECHANISM OF ACTION - Vaccine; gene therapy. Groups of mice were immunized either with the polypeptide (BASB118) adsorbed onto AlPO4 (10 micro g BASB118 onto 100 micro g of AlPO4), with a killed whole cell (kwc) preparation of M. catarrhalis strain American type Culture Collection (ATCC) 43617 adsorbed onto AlPO4, or with 100 micro g AlPO4 without antigen. The mice were challenged with 5 multiply 105 colony forming units (CFU) of live M. catarrhalis strain ATCC 43617 bacteria. The log10 weighted mean number of CFU/lung and the standard deviation 4 hours after challenge was calculated for each group. Sham immunized mice had 5.66 (+/-0.18) log10 CFU/lungs 4 hours after challenge. The kwc preparation induced significant lung clearance as compared to the control group (1.3 log difference). BASB118 vaccine induced a 0.43 log difference in lung clearance, which was significantly different from the control.

USE - A composition comprising an immunologic amount of the new polypeptide or polynucleotide encoding it, is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection (all claimed). The polypeptide may also be used as a prophylactic agent of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptides or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the new polypeptide or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging diseases, and determining the response of an infectious organism to drugs. Dwg.0/1

L13 ANSWER 21 OF 37 WPIDS (C) 2002 THOMSON DERWENT

WPIDS 2001-159870 [16] ACCESSION NUMBER:

N2001-116480 DOC. NO. NON-CPI: C2001-047622

DOC. NO. CPI: New BASB123 polypeptides and TITLE:

polynucleotides from Moraxella catarrhalis strain American type Culture Collection 43617, for use as therapeutic agents or vaccines against bacterial

infections, e.g. otitis media or pneumonia.

DERWENT CLASS: B04 D16 S03

THONNARD, J INVENTOR(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS PATENT ASSIGNEE(S):

COUNTRY COUNT: 94

PATENT INFORMATION:

PATENT NO KIND DATE LA PG: WEEK 79

WO 2001009333 A2 20010208 (200116)* EN RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

> 308-4994 Shears Searcher

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

AU 2000069880 A 20010219 (200129)

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 20010093 AU 20000698	33 A2	WO 2000-EP7296 AU 2000-69880	20000727

FILING DETAILS: .

PATENT NO	KIND	PATENT NO
ALI 20000698	RO A Rased on	WO 200109333

PRIORITY APPLN. INFO: GB 1999-17975 19990730

AN 2001-159870 [16] WPIDS

AB WO 200109333 A UPAB: 20010323

NOVELTY - An isolated polypeptide comprising:

- (a) a sequence comprising one of two 167 amino acid sequences (designated I and II) from Moraxella catarrhalis, given in the specification; or
- (b) an amino acid sequence, which has 85% identity to (I) or (II), over the entire length of (I) or (II), respectively, is new. DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:
- (1) an immunogenic fragment of the new **polypeptide**, in which the immunogenic activity of the fragment is the same as (I) or (II):
- (2) isolated polynucleotides, which encode the new polypeptide, comprising:
 - (i) a nucleotide sequence encoding (a) or (b);
- (ii) a nucleotide sequence that has 85% identity to the nucleotide sequence encoding (I) or (II) over the entire coding region;
- (iii) a 504 base pair (bp) (III) or 501 bp (IV) DNA sequence, given in the specification;
- (iv) a nucleotide sequence that has 85% identity to (III) or (IV) over the entire length of (III) or (IV), respectively;
 - (v) the complements of (i)-(iv); or
- (vi) a nucleotide sequence encoding (I) or (II) obtainable by screening an appropriate library, under stringent conditions, with a labeled probe having (III), (IV), or fragments of (III) or (IV);
- (3) an expression vector or a recombinant live microorganism comprising a polynucleotide of (2);
- (4) a host cell comprising the expression vector of (3), or a subcellular fraction or membrane of the host cell expressing the new polypeptide;
- (5) producing the new polypeptide comprising culturing (4) t produce the polypeptide and recovering it from the culture medium;
 - (6) expressing a polynucleotide of (2) comprising transforming

a host cell with the expression vector of (3) and culturing the host cell for expression of any of the polynucleotides;

(7) vaccine compositions comprising the new polypeptide or polynucleotide of (2), and a pharmaceutical carrier;

(8) an antibody immunospecific for the new

polypeptide or an immunological fragment;

(9) diagnosing a M. catarrhalis infection comprising identifying the new polypeptide or the antibody of (8) present within a biological sample from an animal suspected of having such an infection; and

(10) a therapeutic composition comprising an ${\tt antibody}$ of (8).

ACTIVITY - Antibacterial.

MECHANISM OF ACTION - Vaccine; gene therapy. Clinical details are described but no results are given.

USE - A composition comprising an immunologic amount of the new polypeptide or polynucleotide encoding it, is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection (all claimed). The polypeptides may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptide or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the polypeptide or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging of diseases, and determining the response of an infectious organism to drugs. Dwg.0/2

L13 ANSWER 22 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2001-159869 [16] WPIDS

DOC. NO. NON-CPI:

N2001-116479

DOC. NO. CPI:

C2001-047621

TITLE:

New BASB115 polypeptide from Moraxella

catarrhalis strain MC2931 (ATCC 43617), useful as a therapeutic agent or vaccine against bacterial (especially M. catarrhalis) infections, e.g. otitis

media or pneumonia.

DERWENT CLASS:

B04 D16 S03

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

95

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001009332 A2 20010208 (200116)* EN 72

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000068323 A 20010219 (200129) EP 1204752 A2 20020515 (200239) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	API	PLICATION	DATE
WO 2001009332 AU 2000068323 EP 1204752	= 1 1	AU EP	2000-EP7294 2000-68323 2000-956339 2000-EP7294	20000727 20000727 20000727 20000727

FILING DETAILS:

	TENT NO K					rent no
	2000068323					200109332
EF	1204752	A2	Based	on .	WO	200109332

PRIORITY APPLN. INFO: GB 1999-18003 19990730

AN 2001-159869 [16] WPIDS

AB WO 200109332 A UPAB: 20010323

NOVELTY - A Moraxella catarrhalis strain MC2931 (ATCC 43617) BASB115 polypeptide of 199 amino acids (I) as defined in the specification, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated **polypeptide** (P1) comprising an amino acid sequence which has at least 85%, preferably 100%, identity to (I) over its entire length;
- (2) an immunogenic fragment (P2) of the **polypeptide**, in which the immunogenic activity of the fragment is substantially the same as (I);
 - (3) an isolated polynucleotide (N1) selected from:
 - (a) a nucleotide sequence encoding (I), P1 or P2;
- (b) an isolated polynucleotide comprising a nucleotide sequence encoding a polypeptide that has at least 85%, preferably 95%, identity to (I) over its entire length, or a nucleotide sequence complementary to the isolated polynucleotide;
- (c) an isolated polynucleotide comprising a nucleotide sequence that has at least 85%, preferably 95%, identity to a nucleotide sequence encoding (I) over the entire coding region, or a nucleotide sequence complementary to the isolated polynucleotide;
- (d) an isolated polynucleotide comprising the 600 basepair (bp) sequence (II) fully defined in the specification;
- (e) an isolated polynucleotide comprising a nucleotide sequence which has at least 85%, preferably 95%, identity to (I) over its entire length, or a nucleotide sequence complementary to the isolated polynucleotide;
- (f) a nucleotide sequence encoding (I) obtainable by screening an appropriate library, under stringent conditions, with a labeled

probe having the sequence of (II) or its fragments;

- (4) an expression vector or a recombinant live microorganism comprising N1;
- (5) a host cell comprising the expression vector of (4), or a subcellular fraction or membrane of the host cell expressing P1;
- (6) a process for producing (I), P1 or P2 by culturing the host cell of (5);
- (7) a process for expressing N1 comprising transforming a host cell with the expression vector of (4) and culturing the host cell;
 - (8) a vaccine compositions comprising (I), P1 or P2 or N1;
 - (9) an antibody immunospecific for (I), P1 or P2;
- (10) a method for diagnosing a M. catarrhalis infection comprising identifying (I), P1 or P2 or the **antibody** of (9) present within a biological sample from an animal suspected of having such an infection; and
- (11) a therapeutic composition for treating humans with M. catarrhalis disease, comprising at least one **antibody** against (I), P1 or P2.

ACTIVITY - Antibacterial; ophthalmological; antiinflammatory. MECHANISM OF ACTION - Vaccine; gene therapy.

Groups of mice were immunized either with the polypeptide (BASB115) adsorbed onto AlPO4 (10 mu g BASB115 onto 100 mu g of AlPO4), with a killed whole cells (kwc) preparation of M. catarrhalis strain ATCC 43617 adsorbed onto AlPO4, or with 100 mu g AlPO4 without antigen. The mice were challenged with 5 x 105 colony forming units (CFU) of live M. catarrhalis strain ATCC 43617 bacteria. The log10 weighted mean number of CFU/lung and the standard deviation 4 hours after challenge was calculated for each group. Sham immunized mice had 5.66 (+/-0.18) log10 CFU/lungs 4 hours after challenge. The kwc preparation induced significant lung clearance as compared to the control group (1.76 log difference). BASB115 vaccine induced a 0.46 log difference in lung clearance, which was significantly different from the control.

USE - The composition comprising an immunologic amount of the polypeptide or polynucleotide is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection (all claimed). The polypeptides may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptides or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the polypeptides or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging of the disease, determining the response of an infectious organism to drugs. Dwg.0/1

L13 ANSWER 23 OF 37 WPIDS (C) 2002 THCMSON DERWENT ACCESSION NUMBER: 2001-159854 [16] WPIDS

DOC. NO. CPI:

C2001-047606

TITLE:

New BASB114 polypeptides and

polynucleotides from Moraxella catarrhalis strain

ATCC 43617, useful as therapeutic agents or

vaccines against bacterial infections e.g. otitis

media or pneumonia.

DERWENT CLASS:

B04 D16

95

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

UNIKI COUNI:

PATENT INFORMATION:

PATENT NO	KIND DATE	WEEK	LA	PG

WO 2001009179 A1 20010208 (200116) * EN 82

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE

DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG

KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ

PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW.

AU 2000068322 A 20010219 (200129)

EP 1204678 A1 20020515 (200239) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL RO SI

APPLICATION DETAILS:

PATENT NO K	IND	API	PLICATION	DATE
WO 2001009179 AU 2000068322 EP 1204678		AU EP	2000-EP7293 2000-68322 2000-956338 2000-EP7293	20000727 20000727 20000727 20000727

FILING DETAILS:

PATENT NO	KIND	PATENT NO
AU 20000683	22 A Based on	WO 200109179
EP 1204678	Al Based on	WO 200109179

PRIORITY APPLN. INFO: GB 1999-17977 19990730

AN 2001-159854 [16] WPIDS

AB WO 200109179 A UPAB: 20010323

NOVELTY - An isolated BASB114 Moraxella catarrhalis strain American Type Culture Collection No. 43617 polypeptide (I)

comprising one of two fully defined sequences of 169 amino acids (S1/S2) as given in the specification or an amino acid sequence at least 85% identical to S1/S2, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an immunogenic fragment of (I) in which the immunogenic activity is substantially the same as (I);
 - (2) an isolated polynucleotide (II) comprising:
- (a) a (sequence at least 85% identical to a) nucleotide sequence encoding (I);

- (b) a (sequence at least 85% identical to a) fully defined nucleotide sequence of 510 (S3) or 507 (S4) base pairs (bp) as given in the specification;
 - (c) complements of (a) or (b); or
- (d) a nucleotide sequence obtainable by screening an appropriate library under stringent conditions with a labeled probe containing (fragments of) S3 or S4;
- (3) an expression vector or a recombinant live microorganism
 (III) comprising (II);
- (4) a host cell (IV) comprising (III) or a subcellular fraction or membrane of (IV) expressing (I);
- (5) producing (I) comprising culturing (IV) and recovering the produced polypeptide;
- (6) expressing (II) comprising transforming a host cell with (III) and culturing the host cell;
 - (7) vaccine compositions comprising (I) or (II);
- (8) an **antibody** (V) immunospecific for (I) or its immunological fragment; and
- (9) diagnosing a M. catarrhalis infection comprising identifying (I) or (V) present within a biological sample from an animal suspected of having such an infection.

ACTIVITY - Antibacterial.

MECHANISM OF ACTION - Vaccine; gene therapy.

Groups of mice were immunized either with the polypeptide (BASB114) adsorbed onto AlPO4 (undefined) (10 micro g BASB114 onto 100 micro g of AlPO4), with a killed whole cells (kwc) preparation of M. catarrhalis strain ATCC 43617 adsorbed onto AlPO4, or with 100 micro g AlPO4 without antigen. The mice were challenged with 5 multiply 10 to the power of 5 cell forming units (CFU) of live M. catarrhalis strain ATCC 43617 bacteria. The log 10 weighted mean number of CFU/lung and the standard deviation 4 hours after challenge were calculated for each group. Sham immunized mice had 5.4 (+/-0.2) log 10 CFU/lungs 4 hours after challenge. The kwc preparation induced significant lung clearance as compared to the control group (1.6 log difference). BASB114 vaccine induced a 1.45 log difference in lung clearance, which was significantly different from the control.

USE - The composition comprising an immunologic amount of (I) or (II) is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection (claimed). (I) may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. (II) are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderly patients, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. (I) or (II) may also be employed as research reagents and materials for discovering treatments of and diagnostics for human diseases. In particular, (I) or (II) are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging of the disease, determining the response of an infectious organism to drugs. Dwg.0/4

ACCESSION NUMBER:

2001-112459 [12] WPIDS

DOC. NO. NON-CPI: DOC. NO. CPI:

N2001-082527 C2001-033488

TITLE:

Novel BASB110 polypeptides of Moraxella

catarrhalis, useful as a vaccine for treating

Moraxella catarrhalis infections.

DERWENT CLASS:

B04 D16 S03

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

95

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001000838 A1 20010104 (200112)* EN 88

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE

DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ

PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

AU 2000059779 A 20010131 (200124)

EP 1196589 A1 20020417 (200233) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK

NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	AP	PLICATION	DATE
WO 2001000838 AU 2000059779 EP 1196589		AU EP	2000-EP5854 2000-59779 2000-945812 2000-EP5854	20000623 20000623 20000623 20000623

FILING DETAILS:

11112111 110	KIND		TENT NO
AU 200005977	79 A Based o	n WO	200100838 200100838

PRIORITY APPLN. INFO: GB 1999-15031

19990625

2001-112459 [12] WPIDS ΑN

WO 200100838 A UPAB: 20010302 AB

> NOVELTY - Isolated BASB110 polypeptides (I) of Moraxella catarrhalis, are new. The BASB110 polypeptide has the 322 (P1) or another 322 (P2) amino acid sequence defined in the specification.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated polypeptide (Ia) comprising an amino acid sequence which is at least 85%, preferably 95%, most preferably 100%, identical to the sequence, over its entire length, of P1 or P2;
- (2) an immunogenic fragment (Ib) of (I) or (Ia), where the activity of the fragment is substantially the same as P1 or P2;

Shears 308-4994 Searcher :

- (3) an isolated polynucleotide (II) encoding (I), (Ia) or (Ib);
- (4) an isolated polynucleotide (IIa) comprising a sequence encoding (Ia) or its complementary sequence;
- (5) an isolated polynucleotide (IIb) comprising a sequence having at least 85%, preferably 95%, most preferably 100% identity to a sequence encoding P1 or P2 over the entire coding region, or a nucleotide sequence complementary to the isolated polynucleotide;
- (6) an isolated polynucleotide (IIc) comprising a sequence having at least 85%, preferably 95%, most preferably 100% identical to the 969 (N1) or 966 (N2) nucleotides fully defined in the specification, or its complement;
- (7) an isolated polynucleotide (IId) comprising a sequence encoding P1 or P2, obtainable by screening an appropriate library under stringent hybridization conditions with labeled probe having the sequence of N1 or N2;
- (8) an expression vector (III) of a recombinant live microorganism, comprising (II), (IIa), (IIb), (IIc) or (IId);
- (9) a host cell (IV) comprising (III), or a subcellular fraction or membrane of (IV) expressing (Ia);
- (10) a process for producing (I), (Ia) or (Ib) comprising
 culturing (IV);
- (11) a process for expressing (II), (IIa), (IIb), (IIc) or (IId), comprising transforming (IV) with (III) and culturing transformed (IV) under conditions sufficient for its expression;
- (12) a vaccine composition (V) comprising (I), (Ia) or (Ib), or
 (II), (IIa), (IIb), (IIc) or (IId);
- (13) an **antibody** (Ab1) immunospecific for (I), (Ia) or (Ib); and
- (14) a method for diagnosing Moraxella catarrhalis infection, by identifying (I)-(Ib) or Ab1 present within a biological sample from an animal suspected of having such an infection.

ACTIVITY - Antibacterial.

MECHANISM OF ACTION - Vaccine.

Groups of mice are immunized with BASB110 vaccine. After the booster, the mice were challenged by instillation of bacterial suspension into the nostril under anaesthesia. Mice were killed between 30 minutes and 24 hours after challenge and the lungs were removed aseptically and homogenized individually. The log 10 weighted mean number of colony forming units (CFU)/lung was determined by counting the colonies grown on agar plates after plating of dilutions of the homogenate. The arithmetic mean of the log 10 weighted mean number of CFU/lung and the standard deviations were calculated for each group. Results were not given in the specification.

USE - The vaccine is useful for preparing a medicament for use in generating immune response in an animal (claimed). Abl is useful for treating humans suffering from Moraxella catarrhalis disease (claimed).

Polynucleotides encoding the BASB110 polypeptides have utility in diagnosis of the stage and type of infection, and also for therapeutic or prophylactic purposes, in particular genetic immunization.

Dwg.0/3

L13 ANSWER 25 OF 37 WPIDS (C) 2002 THOMSON DERWENT ACCESSION NUMBER: 2001-112458 [12] WPIDS

DOC. NO. NON-CPI: N2001-082526
DOC. NO. CPI: C2001-033487

TITLE:

New BASB113 polypeptide isolated from

Moraxella catarrhalis bacterium, useful for

diagnosing and producing vaccines against bacterial

infections such as otitis media and pneumonia.

DERWENT CLASS:

B04 D16 S03 THONNARD, J

INVENTOR(S):
PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

OUNTRY COUNT:

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

95

WO 2001000836 A1 20010104 (200112)* EN 86

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE

DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG

KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ

PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

AU 2000059778 A 20010131 (200124)

EP 1196588 A1 20020417 (200233) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	APPLICATION	DATE
WO 2001000836 AU 2000059778 EP 1196588		WO 2000-EP5851 AU 2000-59778 EP 2000-945811 WO 2000-EP5851	20000623 20000623 20000623 20000623

FILING DETAILS:

PAT	ENT NO	KIND				TENT NO
ΑU	200005977	8 A	Based	on	WO	200100836
ΕP	1196588	A1	Based	on	WO	200100836

PRIORITY APPLN. INFO: GB 1999-15044

19990625

AN 2001-112458 [12] WPIDS

AB WO 200100836 A UPAB: 20010302

NOVELTY - An isolated **polypeptide** (I) comprising an amino acid sequence which has 85% identity to the Moraxella catarrhalis BASB113 **polypeptide** sequence of 224 (S2) or 224 (S4) amino acids respectively as given in the specification, or has a sequence of (S2) or (S4), is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an immunogenic fragment (II) of (I) which has the same immunogenic activity as (I);
- (2) an isolated polynucleotide (III), or its complementary nucleotide sequence comprising a nucleotide sequence:
- (i) encoding a polypeptide that has 85% identity over the entire length of (S2) or (S4);
 - (ii) that has 85% identity over the entire length of the

nucleotide sequence encoding region which encodes (S2) or (S4);

- (iii) which has 85% identity over the entire length of a fully defined nucleotide sequence of 675 (S1) or 672 (S3) base pairs as given in the specification; and
- (iv) comprising a nucleotide sequencing encoding (I) obtainable by screening an appropriate library under stringent hybridization conditions with a labeled probe with the sequence of (S1) or (S3);
- (3) an expression vector (IV), or a recombinant live microorganism comprising (III);
- (4) a host cell (V) comprising (IV), or a subcellular fraction or membrane of the host cell expressing (I);
- (5) production of (I) comprising culturing (V) and recovering the produced polypeptide;
- (6) expressing (III) involves transforming (V) with (IV) which contains any one of the polynucleotides given above and culturing (V) under suitable conditions to express the polynucleotides;
 - (7) a vaccine composition which comprises (I) or (II);
 - (8) a vaccine composition which comprises (III);
- (9) an antibody (Ab) immunospecific for (I) or (II); and
- (10) therapeutic compositions comprising an **antibody** directed against (I) useful in treating humans with Moraxella catarrhalis.

ACTIVITY - Anti-inflammatory; auditory; antibacterial.

MECHANISM OF ACTION - Gene therapy; vaccine. Details of test
are given but no results are stated.

USE - (I), (II) and (III) are useful for preparing a medicament useful for generating an immune response in an animal. (I) is also useful as diagnostic reagent for Moraxella catarrhalis which involves identifying (I) or an antibody against (I) present within the biological sample from an animal suspected of having such an infection (claimed). The polynucleotides may be used as hybridization probes for RNA, cDNA and genomic DNA to isolate full-length cDNAs and genomic clones encoding BASB113 and to isolate cDNA and genomic clones of other genes that have high sequence identity to BASB113 gene. The polynucleotides and polypeptides are used as research reagents and materials for discovery of treatments of and diagnostics for human diseases. The polynucleotides derived from (S1) or (S3) is used as PCR (polymerase chain reaction) primers. The polynucleotide sequences can be used in the discovery and development of antibacterial compounds. The encoded protein can be used as target for the screening of antibacterial drugs. Additionally, the polynucleotide sequences encoding the amino terminal regions of the encoded protein or Shine-Dalgarno or other translation facilitating sequences of the respective mRNA can be used to construct antisense sequences to control the expression of the coding sequence of interest. The polypeptides and polynucleotides are used to block the initial physical interaction between a gram negative and/or gram. positive bacteria to mammalian, host thus preventing the sequelae of infection. The polynucleotides encoding certain non-variable regions of bacterial cell surface protein are used in polynucleotide constructs which are useful for genetic immunization experiments in animal models of infection with Moraxella catarrhalis to identify protein groups able to provoke a prophylactic or therapeutic immune response. The vaccine comprising (I), (II) or (III) is useful for treating Moraxella catarrhalis infections such as sinusitis, nosocomial infections, otitis media and pneumonia.

(II) is also used for the rapeutic or prophylactic purposes especially genetic immunization. Dwg.0/3

L13 ANSWER 26 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2001-112457 [12]

DOC. NO. NON-CPI: N2001-082525 DOC. NO. CPI: C2001-033486

TITLE: Novel BASB112 polypeptides of Moraxella

catarrhalis, useful as a vaccine for treating

WPIDS

Moraxella catarrhalis infections.

DERWENT CLASS: B04 D16 S03 INVENTOR(S): THONNARD, J

PATENT ASSIGNEE(S): (SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT: 95

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001000835 A1 20010104 (200112)* EN 81

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW AU 2000061519 A 20010131 (200124)

EP 1196591 A1 20020417 (200233) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	AP	PLICATION	DATE
WO 2001000835 AU 2000061519 EP 1196591		AU EP	2000-EP5849 2000-61519 2000-947873 2000-EP5849	20000623 20000623 20000623 20000623

FILING DETAILS:

PA	TENT NO	KIND			PAI	ENT NO
AU	200006151	9. A	Based	on	WO	200100835
	1196591		Based		WO	200100835

PRIORITY APPLN. INFO: GB 1999-14870 19990625

AN 2001-112457 [12] WPIDS

AB WO 200100835 A UPAB: 20010302

NOVELTY - Isolated BASB112 polypeptides (I) of Moraxella catarrhalis, are new. The BASB112 polypeptide has the 122 (P1) or another 122 (P2) amino acid sequence defined in the specification.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

(1) an isolated polypeptide (Ia) comprising an amino

acid sequence which is at least 85%, preferably 95%, most preferably 100%, identical to the sequence, over its entire length, of P1 or P2;

- (2) an immunogenic fragment (Ib) of (I) or (Ia), where the activity of the fragment is substantially the same as P1 or P2;
 - (3) an isolated polynucleotide (II) encoding (I), (Ia) or (Ib);
- (4) an isolated polynucleotide (IIa) comprising a sequence encoding (Ia) or its complementary sequence
- (5) an isolated polynucleotide (IIb) comprising a sequence having at least 85%, preferably 95%, most preferably 100% identity to a sequence encoding P1 or P2 over the entire coding region, or a nucleotide sequence complementary to the isolated polynucleotide;
- (6) an isolated polynucleotide (IIc) comprising a sequence having at least 85%, preferably 95%, most preferably 100% identical to the 369 (N1) or 366 (N2) nucleotides fully defined in the specification, or its complement;
- (7) an isolated polynucleotide (IId) comprising a sequence encoding P1 or P2, obtainable by screening an appropriate library under stringent hybridization conditions with labeled probe having the sequence of N1 or N2;
- (8) an expression vector (III) of a recombinant live microorganism, comprising (II), (IIa), (IIb), (IIc) or (IId);
- (9) a host cell (IV) comprising (III), or a subcellular fraction or membrane of (IV) expressing (Ia);
- (10) a process for producing (I), (Ia) or (Ib) comprising culturing (IV)
- (11) a process for expressing (II), (IIa), (IIb), (IIc) or (IId), comprising transforming (IV) with (III) and culturing transformed (IV) under conditions sufficient for its expression;
- (12) a vaccine composition (V) comprising (I), (Ia) or (Ib), or (II), (IIa), (IIb), (IIc) or (IId);
- (13) an antibody (Ab1) immunospecific for (I), (Ia)
 or (Ib); and
- (14) a method for diagnosing Moraxella catarrhalis infection, by identifying (I)-(Ib) or Abl present within a biological sample from an animal suspected of having such an infection.

ACTIVITY - Antibacterial.

MECHANISM OF ACTION - Vaccine.

Groups of mice are immunized with BASB112 vaccine. After the booster, the mice were challenged by instillation of bacterial suspension into the nostril under anaesthesia. Mice were killed between 30 minutes and 24 hours after challenge and the lungs were removed aseptically and homogenized individually. The log 10 weighted mean number of colony forming units (CFU)/lung was determined by counting the colonies grown on agar plates after plating of dilutions of the homogenate. The arithmetic mean of the log 10 weighted mean number of CFU/lung and the standard deviations were calculated for each group. Results were not given in the specification.

USE - The vaccine is useful for preparing a medicament for use in generating immune response in an animal (claimed). Abl is useful for treating humans suffering from Moraxella catarrhalis disease (claimed).

Polynucleotides encoding the BASB112 **polypeptides** have utility in diagnosis of the stage and type of infection, and also for therapeutic or prophylactic purposes, in particular genetic immunization.

Dwg.0/3

WPIDS (C) 2002 THOMSON DERWENT L13 ANSWER 27 OF 37

2001-025166 [03] WPIDS ACCESSION NUMBER:

N2001-019583 DOC. NO. NON-CPI: C2001-007779 DOC. NO. CPI:

New BASB103-108 polypeptides isolated TITLE:

from Moraxella catarrhalis bacterium, useful for diagnosing and producing vaccines against bacterial

infections such as otitis media and pneumonia.

B04 D16 S03 DERWENT CLASS: THONNARD, J INVENTOR(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS PATENT ASSIGNEE(S):

COUNTRY COUNT: 94

PATENT INFORMATION:

WEEK LA PG PATENT NO KIND DATE

WO 2000071724 A2 20001130 (200103)* EN 79 RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK

DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP

KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU

ZA ZW

AU 2000045673 A 20001212 (200115)

A2 20020313 (200225) ΕN EP 1185658

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	APPLICATION	DATE .
WO 2000071724 AU 2000045673 EP 1185658		WO 2000-EP4618 AU 2000-45673 EP 2000-927226 WO 2000-EP4618	20000518 20000518 20000518 20000518

FILING DETAILS:

PAT	rent no e	IND			PA'	TENT NO
ΑU	2000045673	Α	Based	on	WO	200071724
EΡ	1185658	A2	Based	on	WO	200071724

PRIORITY APPLN. INFO: GB 1999-13354 19990608; GB 1999-12038 19990524; GB 1999-12040 19990524; GB 19990601; GB 1999-12705 1999-12674

19990601; GB 1999-12838

AN. 2001-025166 [03] WPIDS

WO 200071724 A UPAB: 20010116 AΒ

> NOVELTY - An isolated polypeptide (I) comprising an amino acid sequence which is at least 85% identical to the Moraxella catarrhalis BASB103-BASB108 polypeptides fully defined sequence of 252 (S2), 650 (S4), 405 (S6), 410 (S8), 818 (S10) or 913

(S12) amino acids as given in the specification, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for

Shears 308-4994 Searcher :

the following:

- (1) an immunogenic fragment (II) of (I) which has the same immunogenic activity as (I);
- (2) an isolated polynucleotide (III), or its complementary nucleotide sequence comprising a nucleotide sequence:
 - (a) encoding (I);
- (b) that is 85% identical over the entire sequence which encodes (S2), (S4), (S6), (S8), (S10) or (S12);
- encodes (S2), (S4), (S6), (S8), (S10) or (S12);
 (c) that is 85% identical to a fully defined nucleotide sequence of 759 (S1), 1953 (S3), 1218 (S5), 1233 (S7), 2457 (S9) or 2742 (S11) base pairs as given in the specification; and
- (d) comprising a nucleotide sequencing encoding (I) obtainable by screening an appropriate library under stringent hybridization conditions with a labeled probe having the sequence of (S1), (S3), (S5), (S7), (S9) or (S11);
- (3) an expression vector (IV) or a recombinant live microorganism comprising (III);
- (4) a host cell (V) comprising (IV), or a subcellular fraction or membrane of the host cell expressing (I);
 - (5) preparation of (I);
- (6) expressing (III) involves transforming (V) with (IV) and culturing (V) under suitable conditions to express the polynucleotides;
 - (7) a vaccine composition which comprises (I), (II) or (III);
- (8) an antibody (Ab) immunospecific for (I) or (II); and
- (9) therapeutic compositions comprising an Ab directed against (I).

ACTIVITY - Anti-inflammatory; auditory. No supporting data given.

MECHANISM OF ACTION - Gene therapy; vaccine.

USE - The therapeutic composition comprising (I), an immunogenic fragment (II) of (I) or a polynucleotide (III) encoding (I) is useful for the preparation of a medicament for generating an immune response in an animal. (I) is also useful as a diagnostic reagent for Moraxella catarrhalis which involves identifying (I) or an antibody against (I) present within the biological sample from an animal suspected of having such an infection (claimed). The polynucleotides may be used as hybridization probes for RNA, cDNA and genomic DNA to isolate full-length cDNAs and genomic clones encoding BASB103-108 and to isolate cDNA and genomic clones of other genes that have high sequence identity to BASB103-108 gene. The polynucleotides and polypeptides are used as research reagents and materials for discovery of treatments of and diagnostics for human diseases. The polynucleotides derived from (S1), (S3), (S5), (S7), (S9) or (S11) are used as polymerase chain reaction (PCR) primers. The polynucleotide sequences can be used in the discovery and development of antibacterial compounds. The encoded protein can be used as target for the screening of antibacterial drugs. Additionally, the polynucleotide sequences encoding the amino terminal regions of the encoded protein or Shine-Dalgarno or other translation facilitating sequences of the respective mRNA can be used to construct antisense sequences to control the expression of the coding sequence of interest. The polypeptides and polynucleotides are used to block the initial physical interaction between a gram negative and/or gram positive bacteria to mammalian host thus preventing the sequelae of infection. The polynucleotides encoding certain

non-variable regions of bacterial cell surface **protein** are used in polynucleotide constructs which are useful for genetic immunization experiments in animal models of infection with M.catarrhalis to identify **protein** groups able to provoke a prophylactic or therapeutic immune response. The vaccine comprising (I), (II) or (III) is useful for treating Moraxella catarrhalis infections such as sinusitis, nosocomial infections, otitis media and pneumonia. (II) is also used for therapeutic or prophylactic purposes especially genetic immunization.

Dwg.0/0

L13 ANSWER 28 OF 37

WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2000-587296 [55] WPIDS

DOC. NO. CPI:

C2000-175126

TITLE:

New BASB081 polypeptides from Moraxella catarrhalis and polynucleotides encoding the polypeptides used for treating infections, or as a vaccine for preventing infections, especially those caused by M. catarrhalis.

DERWENT CLASS:

B04 D16

91

INVENTOR(S):

RUELLE, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

PATENT INFORMATION:

PATENT NO KIND DATE W	WEEK LA	PG
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WO 2000052042 A1 20000908 (200055)* EN 97

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW NL OA PT SD SE SL SZ TZ UG ZW
W: AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK DM
EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ

LC LK LR LS LT LU LV MA MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000029136 A 20000921 (200065) EP 1163265 A1 20011219 (200206)

.163265 A1 20011219 (200206) EN R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

APPLICATION DETAILS:

PATENT NO KIND	APPLICATION	DATE
WO 2000052042 A1 AU 2000029136 A EP 1163265 A1	WO 2000-EP1468 AU 2000-29136 EP 2000-907603 WO 2000-EP1468	20000223 20000223 20000223 20000223

FILING DETAILS:

PATENT	NO K	CIND			PAT	ENT	NO
	0029136				WO	2000	52042
EP 116	3265	A1	Based	on	WC	2000	52042

PRIORITY APPLN. INFO: GB 1999-4559

19990226

AN 2000-587296 [55] WPIDS

AB WO 200052042 A UPAB: 20001102

NOVELTY - New isolated BASB081 polypeptides comprising a

sequence of 919 amino acids (Ia), 889 amino acids (Ib), both given in the specification, or a sequence with 85 % identity (Ic) to (Ia) or (Ib), is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an immunogenic fragment of the new **polypeptide** in which the immunogenic activity of the fragment is substantially the same as (Ia) or (Ib);
- (2) polynucleotides with DNA sequences comprising 2760 bp (IIa), 2670 bp (IIb), or a sequence with at least 85 % identity to (Ia) or (IIb) that encode (Ia) - (Ic), respectively;
- (3) an expression vector or a recombinant live microorganism comprising the isolated polynucleotides;
- (4) a host cell comprising the expression vector, a subcellular fraction or a membrane of the host cell expressing the isolated polypeptide comprising an amino acid sequence having at least 85 % identity to (Ia) or (Ib);
- (5) producing the **polypeptides** comprising culturing the host cell for the production of the **polypeptide**, and recovering the **polypeptide** from the culture medium;
- (6) expressing the polynucleotides comprising transforming a host cell with the expression vector, and culturing the host cell for expression of any one of the polynucleotides;
- (7) vaccine compositions comprising any of the polypeptides or any of the polynucleotides;
- (8) an antibody immunospecific for the polypeptide or the immunological fragment;
- (9) diagnosing a Moraxella catarrhalis infection, by identifying any of the polypeptides, or an antibody that is immunospecific for the polypeptide, present within a biological sample from an animal suspected of having such an infection; and
- (10) a therapeutic composition for treating humans with M. catarrhalis disease comprising an **antibody** directed against any of the **polypeptides**.

ACTIVITY - Anti-bacterial; immunostimulant; antiinflammatory. No biological data is given.

MECHANISM OF ACTION - Vaccine. No biological data is given.

USE - Compositions comprising any of the **polypeptides** or polynucleotides encoding them are useful in preparing a medicament for generating an immune response in an animal (claimed). The BASB081 polynucleotides and **polypeptides** are useful in preventing or treating bacterial infections, e.g. otitis media in infants and children, pneumonia in elderlies, sinusitis, nosocomial infections, chronic otitis media, auditive nerve damage, upper respiratory tract infection, or inflammation of the middle ear. The BASB081 polynucleotides and **polypeptides** are also useful as diagnostic reagents for diagnosing infections caused by bacteria, especially M. catarrhalis.

L13 ANSWER 29 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2000-271440 [23] WPIDS

DOC. NO. NON-CPI: N2000-203227 DOC. NO. CPI: C2000-082932

TITLE: Novel BASB034 polynucleotides and polypeptides from Moraxella catarrhalis used to prepare vaccines against bacterial

infections.

DERWENT CLASS: INVENTOR(S):

B04 D16 S03

PATENT ASSIGNEE(S):

RUELLE, J (SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

PATENT INFORMATION:

MC
DM
KZ
SD
MK
E

NL PT RO SE SI CZ 2001000927 A3 20010815 (200157) KR 2001085794 A 20010907 (200218)

HU 2001003945 A2 20020228 (200223)

CN 1326509 A 20011212 (200225)

APPLICATION DETAILS:

PATENT NO K	IND	APPLICATION	DATE
WO 2000015802		WO 1999-EP6781	19990914 19990914
AU 9958632 NO 2001001263	A A	AU 1999-58632 WO 1999-EP6781	19990914
BR 9914492	A	NO 2001-1263 BR 1999-14492	20010313 19990914
EP 1114160 .	A1	WO 1999-EP6781. EP 1999-946171	19990914 19990914
CZ 2001000927	A3	WO 1999-EP6781 WO 1999-EP6781	19990914 19990914
KR 2001085794		CZ 2001-927 KR 2001-703287	19990914 20010314
HU 2001003945		WO 1999-EP6781	19990914 19990914
CN 1326509	Α	HU 2001-3945 CN 1999-813243	19990914

FILING DETAILS:

PATENT NO K	IND	PATENT NO
AU 9958632 BR 9914492 EP 1114160 CZ 2001000927 HU 2001003945		WO 200015802 WO 200015802 WO 200015802 WO 200015802 WO 200015802

PRIORITY APPLN. INFO: GB 1998-20002

N 2000-271440 [23] WPIDS

19980914

Searcher :

Shears

308-4994

AB WO 200015802 A UPAB: 20000516
NOVELTY - Isolated BASB034 polypeptides from Moraxella catarrhalis are new.

DETAILED DESCRIPTION - An isolated BASB034 polypeptide (I) is new, and comprises an amino acid sequence which has at least 85% or 95% identity to, or is, one of the four fully defined 442 amino acid sequences given in the specification ((Ia)-(Id)).

INDEPENDENT CLAIMS are also included for the following:

- (1) an immunogenic fragment of (I) in which the immunogenic activity is substantially the same as (Ia)-(Id);
- (2) an isolated polynucleotide encoding (I), or a complementary nucleotide;
- (3) an isolated polynucleotide which has at least 85% identity to a nucleotide encoding (I), or a complementary nucleotide;
- (4) an isolated polynucleotide (II) which comprises a sequence which has at least 85% or 95% identity to over the entire length of, or is, one of the four fully defined 1329 base pair (bp) sequences given in the specification, or its complement;
- (5) an isolated polynucleotide encoding (Ia)-(Id), obtainable by screening an appropriate library under stringent hybridization conditions with a labeled probe having the sequence of (II), or its fragment;
- (6) an expression vector or recombinant live microorganism comprising (II), or the polynucleotides of (2), (3), and (5);
- (7) a host cell comprising the expression vector of (6), or a subcellular fraction of that cell expressing (I);
- (8) producing (I), comprising culturing the host cell of (7) under conditions sufficient for the production of the polypeptide, and recovering the polypeptide from the culture medium;
- (9) expressing (II) or the polynucleotides of (2), (3) or (5), comprising transforming a host cell with a vector comprising at least one of these polynucleotides, and culturing the cell under conditions sufficient for expression of the polynucleotide;
- (10) a vaccine composition comprising an effective amount of
- (I), (II) or the polynucleotides of (2), (3) or (5);;
- (11) an **antibody** immunospecific for (I), or the fragment of (1);
- (12) diagnosing a Moraxella infection, comprising identifying (I), or an **antibody** that is immunospecific for (I), present within a biological sample from an animal suspected of having such an infection;
- (13) use of a composition comprising an immunologically effective amount of (I) or (II) or the polynucleotides of (2), (3) or (5) in the preparation of a medicament for use in generating an immune response in an animal; and
- (14) a therapeutic composition useful in treating humans with M. catarrhalis, comprising at least one **antibody** directed against (I) and a pharmaceutically acceptable carrier.

ACTIVITY - None given.

MECHANISM OF ACTION - None given.

USE - The polynucleotides and polypeptides may be employed as research reagents and material for the discovery of treatments and diagnostics for diseases, particularly human diseases. They are particularly used to diagnose and treat M. catarrhalis infections (claimed). They can be used for diagnosis of disease, staging of disease, or determining response of an infectious organism to drugs. The polynucleotides may be used as a

source for hybridization probes, and for screening of genetic mutations, serotype, organism or strain identification, identification of mutations in BASB034 sequences, and as components of arrays which are useful for diagnostic and prognostic purposes. The polypeptides can be used to produce antibodies The polypeptides can also be used in vaccine formulations, and to identify agonists and antagonists. The polypeptides, antibodies, agonists and antagonists (which are bacteriostatic) are used for the treatment and prevention of diseases such as otitis media in infants and children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, and chronic otitis media with hearing loss. The polypeptides, agonists and antagonists are also used for screening of antibacterial drugs.

ADVANTAGE - The frequency of Moraxella catarrhalis infections has risen dramatically, and it is no longer common to isolate M. catarrhalis strains that are resistant to standard antibiotics. The BASB034 products of the invention can be used screen for new antibacterial compounds that may target these resistant bacteria. Dwg.0/6

L13 ANSWER 30 OF 37

WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2000-206007 [18] WPIDS

DOC. NO. NON-CPI:

N2000-153181

DOC. NO. CPI:

C2000-063720

TITLE:

New isolated Moraxella catarrhalis BASB023

polypeptides, useful for developing

products for the prevention, treatment and diagnosis of e.g. otitis media, pneumonia,

sinusitis or nosocomial infections.

DERWENT CLASS:

B04 D16 S03 THONNARD, J INVENTOR(S):

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

PATENT INFORMATION:

PATENT	NO	KIND	DATE	WEEK	LA	PG
·						

89

WO 2000009694 A1 20000224 (200018) * EN 98

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW NL OA PT SD SE SL SZ UG ZW

W: AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK DM EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG US UZ VN YU ZA ZW

A 20000306 (200030) AU 9954227

EP 1105492 A1 20010613 (200134) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	AP	PLICATION	DATE
WO 2000009694 AU 9954227 EP 1105492	A1 A A1	AU EF	1999-EP5828 1999-54227 1999-940192 1999-EP5828	19990811 19990811 19990811 19990811

308-4994 Searcher : Shears

FILING DETAILS:

AB

PA	TENT NO	O KINE)		PAT	ENT	NO	
	99542		Based Based		•••		09694 09694	

PRIORITY APPLN. INFO: GB 1998-17824

19980814

2000-206007 [18] WPIDS AN

WO 200009694 A UPAB: 20000412

NOVELTY - An isolated polypeptide comprising an amino acid sequence which has at least 85% identity to an 269 residue amino acid sequence, fully defined in the specification, corresponding to the Moraxella catarrhalis BASB023 polypeptide, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated polypeptide (I) having the 269 residue sequence;
- (2) an isolated polypeptide (II) having a variant 269 residue amino acid sequence, fully defined in the specification;
- (3) an immunogenic fragment of (I) or (II) in which the immunogenic activity of the immunogenic fragment is the same as (I);
- (4) an isolated PN comprising a nucleotide sequence (NS) encoding a polypeptide that has at least 85% identity to (I) over its entire length, or a NS complementary to the isolated PN;
- (5) an isolated PN comprising a NS that has at least 85% identity to a NS encoding a (I) over the entire coding region, or a NS complementary to the isolated PN;
- (6) an isolated PN (III) which comprises a NS which has at least 85% identity to an 810 nucleotide sequence, fully defined in the specification and corresponding to a Moraxella cattarhalis BASB023 polynucleotide, over its entire length, or a NS complementary to the isolated PN;
- (7) an isolated PN comprising a NS encoding (I), obtainable by screening an appropriate library under stringent hybridization conditions with a labeled probe having a sequence (III) or a fragment;
- (8) an isolated PN comprising a variant 810 nucleotide sequence, fully defined in the specification;
- (9) an isolated PN comprising a NS encoding a polypeptide of sequence (II), obtainable by screening an appropriate library under stringent hybridization conditions with a labeled probe having a sequence (III) or a fragment;
- (10) an expression vector or recombinant live microorganism comprising an isolated PN of (4)-(9);
- (11) a host cell comprising an expression vector of (10) or a subcellular fraction or a membrane of the host cell expressing an isolated polypeptide comprising an amino acid sequence that has at least 85% identity to an amino acid sequence (I);
- (12) a process for producing the novel polypeptide, comprising culturing the host cell (11) under expression conditions and recovering the polypeptide;
- (13) a process for expressing a PN of (4)-(9), comprising transforming a host cell with the expression vector comprising on of the PN and culturing under expression conditions;
 - (14) a vaccine composition comprising (I), (II), an immunogenic

fragment of (I) or (II), or a PN of (4)-(9), and a carrier; (15) an antibody immunospecific for (I), (II) or the

immunogenic fragment of (2);

(16) a method of diagnosing a Moraxella infection, comprising identifying (I), (II), the immunogenic fragment of (2) or the antibody of (15) in a biological sample form a suspect animal; and

(17) a therapeutic composition for treating Moraxella catarrhalis disease in humans, comprising at least one antibody of (15), and a carrier.

ACTIVITY - Antibacterial; Auditory; Antiinflammatory. MECHANISM OF ACTION - Vaccine. Polyvalent antisera directed against the BASB023 protein were generated by vaccinating 2 rabbits with the purified recombinant BASB023 protein. Each animal was given a total of 3 immunizations intramuscularly (i.m.) of about 20 mu g BASB023 protein per injection (beginning with complete Freund's adjuvant and followed with incomplete Freund's adjuvant) at approx. 21 day intervals. Animals were bled prior to the first immunization and on days 35 and 57. Anti-BASB023 protein titers were measured by an enzyme linked immunosorbant assay (ELISA) using purified recombinant BASB023 protein (0.5 mu g/well). The titer was defined as the highest dilution at least 0.1 as calculated with the following equation: average OD of 2 test samples of antisera - the average OD of 2 test samples of buffer. The titers after 3 immunizations were above 3000.

USE - The Moraxella catarrhalis can cause diseases such as otitis melia, pneumonia, sinusitis and nosocomial infections. The polypeptides and PNs can be used as vaccines (claimed) to protect against infection, particularly Moraxella catarrhalis infections. The antibodies can be used for treating humans with Moraxella catarrhalis disease (claimed). The detection of the polypeptides or antibodies can be used for diagnosing Moraxella infection (claimed). The products can also be used for detection and drug screening. Dwg.0/6

L13 ANSWER 31 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2000-116286 [10]

DOC. NO. NON-CPI: N2000-088100

DOC. NO. CPI: C2000-035435

TITLE: Novel antigens of Branhamella

catarrhalis used for diagnosis, detection

WPIDS

and in vaccines.

DERWENT CLASS:

B04 D16 S03

INVENTOR(S):

CRIPPS, A W; KYD, J

PATENT ASSIGNEE(S): (COR

(CORT-N) CORTECS UK LTD; (CORT-N) CORTECS OM LTD;

(PROV-N) PROVALIS UK LTD; (CORT-N) CORTECS OM PTY

LTD

COUNTRY COUNT:

87

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 9958563 A2 19991118 (200010) * EN 32

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SL SZ UG ZW

W: AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES

FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG US UZ VN YU ZA ZW AU 9938383 A 19991129 (200018)
EP 1077999 A2 20010228 (200113) EN

R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE NO 2000005670 A 20010110 (200115) CN 1306542 A 20010801 (200172) KR 2001071236 A 20010728 (200208)

JP 2002514657 W 20020521 (200236) 37

APPLICATION DETAILS:

PATENT NO KIND		ENT NO K	IND	APE		DATE	
		9958563 9938383	A2 A		1999-GB1473 1999-38383	19990511 19990511	
		1077999	A2	EP	1999-921008 1999-GB1473	19990511 19990511	
	NO	2000005670	A	WO	1999-GB1473	19990511	
	CN	1306542	А	CN	2000-5670 1999-807588	20001110 19990511	
		2001071236 2002514657			2000-712608 1999-GB1473	20001110 19990511	
			•	JP	2000-548365	19990511	

FILING DETAILS:

PATENT NO		KIND			PAT	PATENT NO	
AU	9938383	A	Based	on	WO	9958563	
EΡ	1077999	A2	Based	on	WO	9958563	
JP	200251465	7 W	Based	on	WO	9958563	

PRIORITY APPLN. INFO: GB 1998-10084 19980511

AN 2000-116286 [10] WPIDS

AB WO 9958563 A UPAB: 20000228

NOVELTY - Novel Branhamella catarrhalis antigens are disclosed.

DETAILED DESCRIPTION - A protein (I) which is a B.

catarrhalis antigen, and which has an apparent molecular weight of about 14-71 kDa (as determined by SDS- PAGE), is new.

INDEPENDENT CLAIMS are also included for the following:

- (1) A homolog or derivative of (I).
- (2) One or more antigenic fragments of (I).
- (3) A nucleic acid (II) molecule comprising:
- (a) a DNA sequence coding for (I), or its RNA equivalent;
- (b) a sequence complementary to (a);
- (c) a sequence which has substantial identity with (a) or (b);
- (d) a sequence which codes for a homolog, derivative or fragment of (I).
 - (4) A vector comprising (II).
- (5) A host cell transformed or transfected with the vector of (4).
- (6) An immunogenic composition which is especially a vaccine, comprising (I), or the **proteins** of (1) or (2).
- (7) The use of (I) or the **proteins** of (1) or (2) in the preparation of an immunogenic composition.
 - (8) An antigen composition, comprising (I) and/or the

proteins of (1) and/or (2), optionally together with at least one other B, catarrhalis antigen, or fragment thereof.

(9) An antibody raised against (I) or the proteins of (1) or (2).

(10) A method for detecting and/or diagnosing B. catarrhalis, comprising bringing into contact the **antibody** of (9), (I), the **proteins** of (1) or (2), or the antigen composition of

(8) with a sample to be tested, and detecting the presence of (I).

(11) The use of (I), the **proteins** of (1) or (2), or the antigen composition of (8) in detecting and/or diagnosing B. catarrhalis.

(12) A kit for use in detecting and/or diagnosing B. catarrhalis, comprising (I), the **proteins** of (1) or (2), the antigen composition of (8) or the **antibody** of (9).

(13) The use of (I), or the **proteins** of (1) or (2) or the immunogenic composition of (8) in medicine, or for inducing an immune response in a subject.

(14) A method for the treatment or prophylaxis of respiratory infection or otitis media in a subject, comprising administering an effective amount of (I), the **proteins** of (1) or (2) or the immunogenic composition of (8).

USE - The antigens can be used to prepare vaccines and immunogenic compositions for the treatment and prophylaxis of Branhamella catarrhalisinfections, respiratory tract infections, and otitis media (claimed). Antibodies against the antigens can be used for diagnosis and purification of the antigens.

ADVANTAGE - A need exists for antigens from Branhamella catarrhalis to provide better and more effective vaccines. This need is met by the antigens of the invention.

Dwg.0/0

L13 ANSWER 32 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2000-062302 [05] WPIDS

DOC. NO. NON-CPI: N2000-048800 DOC. NO. CPI: C2000-017246

TITLE: Novel peptides useful for diagnosis,

prophylaxis and treatment of Moraxella infections such as otitis media, pneumonia, sinusitis etc..

DERWENT CLASS: B04 D16 S03 INVENTOR(S): RUELLE, J

PATENT ASSIGNEE(S): (SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT: 87

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 9958685 A2 19991118 (200005) * EN 87

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SL SZ UG ZW

W: AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG

SI SK SL TJ TM TR TT UA UG US UZ VN YU ZA ZW

AU 9942602 A 19991129 (200018) EP 1078066 A2 20010228 (200113)

R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

EN

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 9958685 AU 9942602 EP 1078066	A2 A · · A2	WO 1999-EP3263 AU 1999-42602 EP 1999-950354 WO 1999-EP3263	19990510 19990510 19990510 19990510

FILING DETAILS:

PATENT NO	KIND	PATENT NO
AU 9942602	A Based on	WO 9958685
EP 1078066	A2 Based on	WO 9958685

PRIORITY APPLN. INFO: GB 1999-9175 19990421; GB 1998-10379

19980513

AN 2000-062302 [05] WPIDS

AB WO 9958685 A UPAB: 20000128

NOVELTY - An isolated polypeptide with the Moraxella catarrhalis BASB028 polypeptide (I) sequence of 1726 amino. acids fully defined in the specification, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated polypeptide (II), comprising an amino acid sequence which has 85% identity to the amino acid sequence of (I);
- (2) an immunogenic fragment (III), of (I) or (II) which has the same immunogenic activity as (I);
- (3) an isolated polynucleotide (IV), comprising a nucleotide sequence encoding (I);
- (4) an isolated polynucleotide (V), or its complementary nucleotide sequence comprising a nucleotide sequence:
- (a) encoding a **polypeptide** that has 85% identity over the entire length of (I);
- (b) that has 85% identity over the entire length of the nucleotide sequence coding region which encodes (I); and
- (c) which has 85% identity over the entire length of a fully defined nucleotide sequence of 5181 base pairs (1) as given in the specification;
- (5) an expression vector (VI), or a recombinant live microorganism comprising (IV) or (V);
- (6) a host cell (VII), or a membrane comprising (VI) which expresses (II);
- (7) preparation of (I), comprising culturing host cells of (6) to produce the **polypeptide**, and recovering it from the culture medium;
- (8) expression of (IV) or (V) which comprises transforming (VII) with (VI) which contains any one of the polynucleotides given above and culturing (VII) under suitable conditions to express the polynucleotides;
 - (9) a vaccine composition which comprises (I) or (II);
 - (10) a vaccine composition which comprises (IV) or (V);
- (11) an antibody (Ab) immunospecific for (I), (II) or (III); and
- (12) diagnosing a Moraxella injection by identifying (I), (II), (III) or an Ab produced against them, present in a biological

MECHANISM OF ACTION - Vaccine The efficacy of BASB028 vaccine was analyzed by enhancement of lung clearance of M.catarrhalis in mice. Groups of 6 BALB/c mice were immunized subcutaneously with 100 mu l of vaccine corresponding to a 10 mu l dose and were boosted 2 weeks later. One week after the booster, the mice were challenged by instillation of 50 mu l of bacterial suspension into the left nostril under anesthesia and 0.8 mg ketamine. Mice were killed 4 hours after challenge and the lungs are removed aseptically an homogenized individually. The log 10 weighted mean number of CFU/lung is determined by counting the colonies grown on Mueller-Hinton agar plates after plating of 20 mu l of 5 serial dilutions of the homogenate. No results of the test were given.

USE - The polynucleotides may be used as hybridization probes for RNA, cDNA and genomic DNA to isolate full-length cDNAs and genomic clones encoding BASB028 and to isolate cDNA and genomic clones of other genes that have high sequence identity to BASB028 gene. The polynucleotides and polypeptides are used as research reagents and materials for discovery of treatments of and diagnostics for human diseases. The polynucleotides derived from (1) are used for PCR to determine whether or not the identified polynucleotides are transcribed in bacteria in infective tissue and so are helpful in the diagnosis of the stage and type of infection, the pathogen has attained. Probes comprising BASB028 nucleotide sequence can be constructed to conduct efficient screening of genetic mutations, serotype, taxonomic classification or identification. Primers with 1-4 nucleotides removed from the 5' and/or 3' end are used for amplifying BASB028 DNA and/or RNA isolated from a sample derived from an individual. The polynucleotides are used as components of high density polynucleotide arrays or grids which are useful for diagnostic and prognostic purposes. The antibodies directed against (I) or (IV) are employed to isolate or to identify clones expressing (I) or (IV) or to purify them. The polynucleotide sequences can be used in the discovery and development of antibacterial compounds. The encoded protein, for expression can be used as target for the screening of antibacterial drugs. Additionally, the polynucleotide sequences encoding the amino terminal regions of the encoded protein or Shine-Delgarno or other translation facilitating sequences of the respective mRNA can be used to construct antisense sequences to control the expression of the coding sequence of interest. The polypeptides and polynucleotides are used to block the initial physical interaction between a gram negative and/or gram positive bacteria to the mammalian host. The polynucleotides encoding certain non-variable regions of bacterial cell surface protein are used in polynucleotide constructs which are useful for genetic immunization experiments in animal models of infection with M.catarrhalis to identify protein epitopes able to provoke a prophylactic or therapeutic immune response. The therapeutic composition comprising an immunologically effective amounts of a polypeptide, (I) or (II); or a polynucleotide, (IV) or (V) is useful in the preparation of a medicament for generating an immune response in an animal. A therapeutic composition comprising an Ab directed against one or two useful for treating humans with M.catarrhalis diseases (claimed) such as sinusitis, otitis media and

nosocomial infections. Dwg.0/1

L13 ANSWER 33 OF 37 WPIDS (C) 2002 THOMSON DERWENT ACCESSION NUMBER: 2000-062301 [05] WPIDS

ACCESSION NUMBER: 2000-062301 [05] DOC. NO. NON-CPI: N2000-048799

DOC. NO. NON-CPI: N2000-048799 DOC. NO. CPI: C2000-017245

TITLE: Novel peptides useful as vaccines for Moraxella infections such as otitis media,

pneumonia, sinusitis etc.,.

DERWENT CLASS: B04 D16 S03

INVENTOR(S): THOHNARD, J; THONNARD, J

PATENT ASSIGNEE(S): (SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT: 87

PATENT INFORMATION:

PAT	TENT	NO	F	KIND	D DA	ATE		WI	EEK]	ĹΑ	PC	3							
WO	995	3684	1	A2	2 19	9991	1118	3 (;	2000	005)	* F	EN	113	3							
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				LT													RO	RU	SD	SE	SG
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	994																				
EP	1078																				
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	737																				
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APPLICATION DETAILS:

PAT	ENT NO K	IND	API	PLICATION	DATE
WO	9958684	A2		1999-EP3257	19990507
ΑU	9941421	Α .	ΑU	1999-41421	19990507
ΕP	1078064	A2	ΕP	1999-924948	19990507
	•		WO	1999-EP3257	19990507
NO	2000005697	A	WO	1999-EP3257	19990507
			NO	2000-5697	20001110
ÇZ	2000004203	A3	WO	1999-EP3257	19990507
			CZ	2000-4203	19990507
ÁU	737196	В	ΑU	1999-41421	19990507
KR	2001043573	A	KR	2000-712705	20001113
CN	1309706	A ·	CN	1999-808554	19990507
HU	2001002853	A2	WO	1999-EP3257	19990507
			HU	2001-2853	19990507
ZA	2000006522	A	ZA	2000-6522	20001110
ВR	9911773	A	BR	1999-11773	19990507

	WO 1999-EP3257	19990507
MX 2000011140 A1	MX 2000-11140	20001113
JP 2002514425 W	WO 1999-EP3257	19990507
	JP 2000-548475	19990507

FILING DETAILS:

PATENT NO KIND	PATENT NO
AU 9941421 A Based on EP 1078064 A2 Based on	WO 9958684 WO 9958684
CZ 2000004203 A3 Based on	WO 9958684
AU 737196 B Previous Publ. Based on	AU 9941421 WO 9958684
HU 2001002853 A2 Based on	WO 9958684
BR 9911773 A Based on	WO 9958684
JP 2002514425 W Based on	WO 9958684

PRIORITY APPLN. INFO: GB 1998-10285 19980513

AN 2000-062301 [05] WPIDS

AB WO 9958684 A UPAB: 20000128

NOVELTY - An isolated polypeptide with Moraxella

catarrhalis BASB020 polypeptide (I),(II),(III),(IV)

sequence of 280 amino acids (aa) as given in the specification, from M.catarrhalis strains MC2931, MC2912, MC2913 and MC2969, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated polypeptide (V), comprising an aa sequence which has 85% identity to the aa sequence of (I), (II), or (IV);
- (2) an immunogenic fragment (VI), of (I),(II),(III),(IV) or
 (V) which has the same immunogenic activity as (I),(II),(III) or
 (IV);
- (3) an isolated polynucleotide (VII), comprising a nucleotide sequence encoding (I), (II), (III) or (IV);
- (4) an isolated polynucleotide (VII), or its complementary nucleotide sequence comprising a nucleotide sequence:
- (a) encoding a polypeptide that has 85% identity overthe entire length of (I), (II), (III) or (IV);
- (b) that has 85% identity over the entire length of the nucleotide sequence coding region which encodes (I),(II),(III) or (IV); and
- (c) which has 85% identity over the entire length of a fully defined nucleotide sequence of 843 base pairs (1,2,3,4) as given in the specification;
- (5) an expression vector (IX), or a recombinant live microorganism comprising (VII) or (VIII);
- (6) a host cell (X), or a membrane comprising (IX) which expresses (V);
 - (7) preparation of (I),(II),(III) or (IV);
- (8) expression of (VII) or (VIII) which comprises transforming (X) with (IX) which contains any one of the polynucleotides given above and culturing (X) under suitable conditions to express the polynucleotides;
- (9) a vaccine composition which comprises (I),(II),(III) or
 (IV) or (V);
 - (10) a vaccine composition which comprises (VII) or (VIII);
 - (11) an antibody (Ab) immunospecific for

(I), (II), (III), (IV), (V) or (VI); and

(12) diagnosing a Moraxella infection by identifying (I),(II),(III), (IV),(V) or (VI) or an Ab produced against them, present in a biological sample obtained from an animal suspected of having such infection.

ACTIVITY - Anti-inflammatory; auditory.

MECHANISM OF ACTION - Vaccine. The efficacy of BASB020 vaccine was analyzed by enhancement of lung clearance of M.catarrhalis in mice. Groups of 6 BALB/c mice were immunized subcutaneously with 100 mu l of vaccine corresponding to a 10 mu l dose and were boosted 2 weeks later. One week after the booster, the mice were challenged by instillation of 50 mu l of bacterial suspension into the left nostril under anesthesia and 0.8 mg ketamine. Mice were killed 4 hours after challenge and the lungs are removed aseptically a homogenized individually. The log 10 weighted mean number of CFU/lung is determined by counting the colonies grown on Mueller-Hinton agar plates after plating of 20 mu l of 5 serial dilutions of the homogenate. BASB020 vaccine induced significant lung clearance as compared to the control (0.62 log difference).

USE - The polynucleotides may be used as hybridization probes for RNA, cDNA and genomic DNA to isolate full-length cDNAs and genomic clones encoding BASB020 and to isolate cDNA and genomic clones of other genes that have high sequence identity to BASB020 gene. The polynucleotides and polypeptides are used as research reagents and materials for discovery of treatments of and diagnostics for human diseases. The polynucleotides derived from (1,2,3,4) are used for PCR to determine whether or not the identified polynucleotides are transcribed in bacteria in infective tissue and so are helpful in the diagnosis of the stage and type of the pathogen has attained. Probes comprising BASB020 infection, nucleotide sequence can be constructed to conduct efficient screening of genetic mutations, serotype, taxonomic classification or identification. Primers with 1-4 nucleotides removed from the 5' and/or 3' end are used for amplifying BASB020 DNA and/or RNA isolated from a sample derived from an individual. The polynucleotides are used as components of high density polynucleotide arrays or grids which are useful for diagnostic and prognostic purposes. The antibodies directed against (I),(II),(III),(IV) or (VII) are employed to isolate or to identify clones expressing (I),(II),(III),(IV) or (VII) or to purify them. The polynucleotide sequences can be used in the discovery and development of antibacterial compounds. The encoded protein , for expression can be used as target for the screening of antibacterial drugs. Additionally, the polynucleotide sequences encoding the amino terminal regions of the encoded protein or Shine-Delgarno or other translation facilitating sequences of the respective mRNA can be used to construct antisense sequences to control the expression of the coding sequence of interest. The polypeptides and polynucleotides are used to block the initial physical interaction between a gram negative and/or gram positive bacteria to the mammalian host. The polynucleotides encoding certain non-variable regions of bacterial cell surface protein are used in polynucleotide constructs which are useful for genetic immunization experiments in animal models of infection with M.catarrhalis to identify protein epitopes able to provoke a prophylactic or therapeutic immune response. The therapeutic composition comprising an immunologically effective amounts of a polypeptide, (I),(II),(III),(IV) or (V); or a

polynucleotide, (VII) or (VIII) is useful in the preparation of a medicament for generating an immune response in an animal. A therapeutic composition comprising an Ab directed against one or two useful for treating humans with M.catarrhalis diseases (claimed) such as sinusitis, otitis media and nosocomial infections. Dwg.0/8

L13 ANSWER 34 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2000-062033 [05] WPIDS

DOC. NO. NON-CPI: N2000-048594 DOC. NO. CPI: C2000-017145

TITLE: New polypeptides from Moraxella

catarrhalis used to treat the infection by this

bacteria.

DERWENT CLASS: B04 D16 S03 INVENTOR(S): RUELLE, J

PATENT ASSIGNEE(S): (SMIK) SMITHKLINE BEECHAM BIOLOGICALS.

COUNTRY COUNT: 87

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 9955871 A1 19991104 (200005)* EN 70 RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW NL OA PT SD SE SL SZ UG ZW

W: AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG

SI SK SL TJ TM TR TT UA UG US UZ VN YU ZA ZW

AU 9940331 A 19991116 (200015) EP 1071784 A1 20010131 (200108) EN

R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE ·
WO 9955871	A1	WO 1999-EP2764	19990420
AU 9940331	. A	AU 1999-40331	19990420
EP 1071784	A1	EP 1999-923457	19990420
		WO 1999-EP2764	19990420

FILING DETAILS:

PATENT NO	·KIND	PATENT NO
AU 9940331	A Based on	WO 9955871
EP 1071784	Al Based on	WO 9955871

PRIORITY APPLN. INFO: GB 1998-8720 19980423

AN 2000-062033 [05] WPIDS AB WO 9955871 A UPAB: 2000

WO 9955871 A UPAB: 20000128 NOVELTY - Polypeptides from Moraxella catarrhalis,

designated BASB011, are new.

DETAILED DESCRIPTION - An isolated polypeptide (P1)

has an amino acid (aa) sequence having at least 85% identity to one of the sequences fully defined in the specification.

INDEPENDENT CLAIMS are also include for the following:

- (1) an immunogenic fragment of P1, where immunogenic activity is substantially the same as P1;
- (2) an isolated polynucleotide comprising a sequence encoding P1, or its complement;
- (3) an isolated polynucleotide comprising a sequence having at least 85 (preferably at least 95)% identity to a sequence encoding P1 or its complement;
- (4) an isolated polynucleotide comprising a nucleotide sequence having at least 85 (preferably at least 95)% identity over its full length to one of the sequences fully defined in the specification;
- (5) an expression vector or recombinant live organism comprising one of the above polynucleotides;
- (6) a host cell comprising the above expression vector, or a membrane of that host cell expressing P1;
- (7) producing P1, comprising culturing the above host cell under production conditions and recovering the polypeptide
- (8) a vaccine comprising P1 or one of the above polynucleotides in combination with at least one other Moraxella catarrhalis antigen;
- (9) diagnosing a Moraxella infection, comprising identifying P1 or an antibody specific for P1 in a biological sample from an animal, and
- (10) a composition for treating humans with Moraxella disease, comprising at least one antibody directed against P1.

USE - The polypeptide is used to generate an immune response in an animal (claimed), particularly against a bacterial infection, e.g. a Moraxella catarrhalis infection. M. catarrhalis is present in 15% of childhood middle ear infections in the US. Molecules of the invention may also be used to prevent adhesion of bacteria to extracellular matrix proteins on indwelling devices or in wounds, to block bacterial adhesion between extracellular matrix proteins and BASB011 proteins that mediate tissue damage, or to block the normal progression of pathogenesis in infections initiated other than by implanting of indwelling devices or by other surgical techniques.

ADVANTAGE - None given

Dwg.0/17

L13 ANSWER 35 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2000-039107 [03] WPIDS

DOC. NO. NON-CPI: N2000-029453

DOC. NO. CPI: C2000-010168

TITLE: Novel BASB010 polynucleotides and

polypeptides from Moraxella catarrhalis
used to prepare vaccines against bacterial

infections. B04 D16 S03

DERWENT CLASS: B04 D16 S03 INVENTOR(S): THONNARD, J

PATENT ASSIGNEE(S): (SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT: 87

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 9958682 A2 19991118 (200003)* EN 100

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SL SZ UG ZW

W: AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG US UZ VN YU ZA ZW

AU 9942600 A 19991129 (200018)

EP 1078065 A2 20010228 (200113) EN

R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 9958682 AU 9942600 EP 1078065	A2 A A2	WO 1999-EP3254 AU 1999-42600 EP 1999-950353	19990507 19990507 19990507
	A2		

FILING DETAILS:

PATENT NO	KIND	PATENT NO
AU 9942600	A Based on	WO 9958682
EP 1078065	A2 Based on	WO 9958682

PRIORITY APPLN. INFO: GB 1999-5308

19990308; GB 1998-10195

19980512

AN 2000-039107 [03] WPIDS

AB WO 9958682 A UPAB: 20000118

NOVELTY - Novel BASB010 polynucleotides and polypeptides from Moraxella catarrhalis are disclosed.

DETAILED DESCRIPTION - An isolated BASB010 **polypeptide** (I) is new, and comprises an amino acid sequence which has at least 85% or 95% identity to, or is, the 391 (Ia), 391 (Ib) or 391 (Ic) amino acid sequences given in the specification.

INDEPENDENT CLAIMS are also included for the following:

- (1) An immunogenic fragment of (I) in which the immunogenic activity is substantially the same as (Ia), (Ib) or (Ic);
- (2) An isolated polynucleotide encoding (I), or a complementary nucleotide;
- (3) An isolated polynucleotide (II) which comprises a sequence which has at least 85% or 95% identity to over the entire length, or is, the 1176 bp (IIa), 1176 bp (IIb) or 1176 bp (IIc) sequence given in the specification, or its complement;
- (4) An isolated polynucleotide encoding (Ia)-(Ic), obtainable by screening an appropriate library under stringent hybridization conditions with a labeled probe having the sequence of (IIa), (IIb), (IIc) or a fragment thereof;
- (5) An expression vector or recombinant live microorganism comprising (II), or the polynucleotides of (2) or (4);
- (6) A host cell comprising the expression vector of (5), or a subcellular fraction of that cell expressing (I);
- (7) A process for producing (I), comprising culturing a host cell under conditions sufficient for the production of the polypeptide, and recovering the polypeptide from the culture medium;
- (8) A process for expressing (II) or the polynucleotides of (2) or (4), comprising transforming a host cell with a vector comprising at least one of these polynucleotides, and culturing the cell under

conditions sufficient for expression of the polynucleotide;

(9) A vaccine composition comprising an effective amount of (I) and a pharmaceutically acceptable carrier;

- (10) A vaccine composition comprising an effective amount of (II) or the polynucleotides of (2) or (4), and a pharmaceutically acceptable carrier;
- (11) An antibody immunospecific for (I), or the fragment of (1);
- (12) A method for diagnosing a M. catarrhalis infection, comprising identifying (I), or an **antibody** that is immunospecific for (I), present within a biological sample from an animal suspected of having such an infection;
- (13) Use of a composition comprising an immunologically effective amount of (I) or (II) or the polynucleotides of (2) or (4) in the preparation of a medicament for use in generating an immune response in an animal; and
- (14) A therapeutic composition useful in treating humans with M. catarrhalis, comprising at least one **antibody** directed against (I) and a pharmaceutically acceptable carrier.

ACTIVITY - Anti-bacterial, immunostimulant.

MECHANISM OF ACTION - Vaccine.

USE - The polynucleotides and polypeptides may be employed as research reagents and material for the discovery of treatments and diagnostics for diseases, particularly human diseases. They can be used for diagnosis of disease, staging of disease, or determining response of an infectious organism to drugs. The polynucleotides may be used as a source for hybridization probes, and for screening of genetic mutations, serotype, organism or strain identification, identification of mutations in BASB013 sequences, and as components of arrays which are useful for diagnostic and prognostic purposes. The polypeptides can be used to produce antibodies. The polypeptides can also be used in vaccine formulations, and to identify agonists and antagonists. The polypeptides, antibodies, agonists and antagonists (which are bacteristatic) are used for the treatment and prevention of diseases such as otitis media in infants and children, pneumonia in the elderly, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in middle ear, auditive nerve damage, delayed speech learning, infection of the upper respiratory tract and inflammation of the middle ear. They are particularly used to diagnose and treat M. catarrhalis infections. The polypeptides, agonists and antagonists are also used for screening of antibacterial drugs.

ADVANTAGE - The frequency of Moraxella catarrhalis infections has risen dramatically, and it is no longer common to isolate M. catarrhalis strains that are resistant to standard antibiotics. The BASB010 products of the invention can be used screen for new antibacterial compounds that may target these resistant bacteria. Dwg.0/4

L13 ANSWER 36 OF 37

WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: CROSS REFERENCE:

2000-038242 [03] WPIDS

DOC. NO. CPI:

1993-093726 [11]; 2000-012250 [01] C2000-009691

TITLE:

Purified Moraxella catarrhalis outer membrane

proteins useful for vaccinating against

chronic otis media, acute maxillary sinusitis and

other bronchopulmonary and lower respiratory tract

infections. B04 D16

DERWENT CLASS:

INVENTOR(S):

HANSEN, E J; HELMINEN, M E; MACIVER, I

PATENT ASSIGNEE(S):

COUNTRY COUNT:

(TEXA) UNIV TEXAS

PATENT INFORMATION:

ENT		 DATE	WEEK	LA	PG
	3826		(200003)*		50

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
US 5993826	A CIP of CIP of	US 1991-745591 WO 1992-US6869 US 1993-25363	19910815 19920814 19930302

FILING DETAILS:

PATENT	NO .	KIND		•	PAT	ENT	NO	
			. – – – –					_
ris 599	3826	Δ	CTP	of	US	5552	2146	

PRIORITY APPLN. INFO: US 1993-25363 19930302; US 1991-745591 19910815; WO 1992-US6869 19920814

AN 2000-038242 [03] WPIDS

CR 1993-093726 [11]; 2000-012250 [01]

AB US 5993826 A UPAB: 20000925

NOVELTY - A purified Moraxella catarrhalis (also called Branhamella catarrhalis and Neisseria catarrhalis) 80 kiloDalton (kD) CopB outer membrane protein (I), is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for:

- (i) an antigen composition (II) prepared by:
- (1) introducing a recombinant expression vector including a DNA segment encoding (I) into a recombinant host cell;
- (2) culturing the host cell under suitable conditions for the expression of (I); and
 - (3) collecting the expressed antigen; and
- (ii) a method (III) for inducing an antibody response to M. catarrhalis 80 kD CopB antigens in an animal, comprising administering (I).

ACTIVITY - Auditory; Respiratory active.

MECHANISM OF ACTION - Vaccine, administration of (I) stimulates an immune response against M. catarrhalis antigens in a patient.

Groups of mice were immunized with the 8B6 monoclonal antibody, specific for the 80 kD outer membrane protein of M. catarrhalis. Control mice were immunized with an irrelevant antibody, 2H11 which is specific for Haemophilus ducreyi. Doses of 150 micrograms were used 18 hours prior to bacterial challenge. 5 Microliter doses of bacterial suspension, containing M. catarrhalis strain 035E, were inoculated into the lungs of the mice. 6 Hours after inoculation, the mice were sacrificed and the number of bacteria remaining in the lungs

was determined. It was found that where the 2H11 antibody was used, 97% of the initial bacterial population remained. However, just 38% remained when the 8B6 antibody was used.

USE - (I) may be used to vaccinate against M. catarrhalis, a pathogen implicating in causing chronic otis media, acute maxillary sinusitis and other bronchopulmonary and lower respiratory tract infections.

Dwg.0/13

L13 ANSWER 37 OF 37 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

1998-377595 [32] WPIDS

DOC. NO. CPI:

C1998-114707

TITLE:

New peptide(s) containing the core epitope of Moraxella catarrhalis Usp proteins - useful in, e.g. vaccines to

prevent or treat M. catarrhalis infection, and

antibodies for passive immunisation.

DERWENT CLASS:

B04 D16

. 82

INVENTOR(S):

AEBI, C; COPE, L D; FISKE, M J; FREDENBURG, R;

HANSEN, E J; MACIVER, I; FREDENBURG, R A

PATENT ASSIGNEE(S):

(TEXA) UNIV TEXAS SYSTEM; (AMCY) AMERICAN CYANAMID

CO; (TEXA) UNIV TEXAS

COUNTRY COUNT:

PATENT INFORMATION:

PA	CENT	ИО		KINI) D?	ATE		WI	EEK			LA	P(
WO	982																				
	RW:	AΤ	BE	CH	DΕ	DK	EΑ	ES	FI	FR	GB	GH	GM	GR	ΙE	ΙT	KE	LS	LU	MC	MW
		NL	OA	PT	SD	SE	SZ	UG	ZW												
	W:	AL	AM	ΑT	UΑ	ΑZ	BA	BB	BG	BR	BY	CA	CH	CN	CU	CZ	DE	DK	EΕ	ES	FI
		GB	GΕ	GH	GM	GW	HU	ID	IL	IS	JP	ΚE	KG	ΚP	KR	ΚZ	LC	LK	LR	LS	LT
		LU	LV	MD	MG	MK	MN	MW	MX	NO	NZ	PL	PT	RO	RU	SD	SE	SG	SI	SK	\mathtt{SL}
		ΤJ	TM	TR	TT	UA	UG	US	UZ	VN	YU	ZW									
ΑU	985	720:	l	Α	19	986	071	7 (:	1998	348))								•		
EΡ	948																				
	R:	AL	AT	BE	CH	DE	DK	ES	FI	FR	GB	GR	ΙE	ΙT	LI	LT	LÜ	r_{Λ}	MC	NL	PT
		RO	SE	SI																	
BR	971	4160)	Α	20	0000	0502	2 (2	2000)33))										
CN	125	1613	L	Α	20	0000	0426	5 (2	2000)36))										
	2000		-		_			•		•											
	200												250)							
US	6310	0190) .	B1	. 20	0013	1030) (2	2003	172))										

APPLICATION DETAILS:

AU 746442

PATENT NO K	IND	APPLICATION	DATE
WO 9828333 AU 9857201	A2 A	WO 1997-US23930 AU 1998-57201	19971219 19971219
EP 948625	A2	EP 1997-953461 WO 1997-US23930	19971219 19971219
BR 9714160	A	BR 1997-14160 WO 1997-US23930	19971219 19971219
CN 1251611	A	CN 1997-180843	19971219
KR 2000057575		WO 1997-US23930 KR 1999-705332	19971219 19990615

20020502 (200238)

JР	2001515467	W		WO	1997-US23930	19971219
				JP	1998-529075	19971219
US	6310190	В1	Provisional	US	1996-33598P	19961220
			Cont of	WO	1997-US23930	19971219
•				US	1999-336447	19990621
ΑU	746442	В		AU	1998-57201	19971219

FILING DETAILS:

PATENT NO K	IND			PATENT NO	
		Based on		WO 9828333	
		Based on Based on		WO 9828333 WO 9828333	
KR 2000057575 JP 2001515467				WO 9828333 WO 9828333	
	В	Previous P	ubl.	AU 9857201	
		Based on		WO 9828333	

PRIORITY APPLN. INFO: US 1996-33598P 19961220; US 1999-336447 19990621

1998-377595 [32] WPIDS

AN

9828333 A UPAB: 19991122 AB

Isolated peptides (I) of 7-60 amino acids (aa) that include the sequence AQQQDQH (S1) are new. Also new are: (1) antiquenic composition or vaccine (A) containing (I) plus buffer or diluent; (2) nucleic acid (II) encoding the UspA1 and A2 antigens of Moraxella catarrhalis

isolates O35E, O46E, TTA24 and TTA37; specific a sequences together with their corresponding coding nucleotide sequences are given in the specification; (3) a method of screening peptides for reactivity with an antibody (Ab) that binds UspA1 or A2; (4) isolated peptides (III) with at least 7 consecutive aa from UspA1 or A2, including residues within the 582-604 or 355-377 aa regions of UspA1 and A2, respectively, of O35E, or analogous regions in other isolates; (5) antigenic construct containing (III) plus buffer or diluent, and (6) antigenic construct containing an isolated 7-60 aa peptide that includes at least 7 aa from UspA1 or A2, acting as a carrier covalently coupled to second antigen.

USE - (A) are used to induce an immune response in mammals against M. catarrhalis ((II) can be used similarly in genetic vaccination) and (I) can be used to treat infections by M. catarrhalis (claimed) (e.g. otitis media, sinusitis, lower respiratory tract infections), and also as immunity enhancers for other bacterial, parasitic or viral antigens, to raise Ab and as immunoassay reagents for detecting specific antibodies. Ab are useful for passive immunisation and as immunoassay reagents. Detection of the epitopic core sequence (i.e. (S1)), by immunoassay or by PCR, is used to diagnose infection (claimed). (II) are also used to produce recombinant proteins and for screening for potential anti-M. catarrhalis agents, while fragments of (II) are useful as diagnostic probes or primers or to isolate variant sequences. (A) are generally administered by subcutaneous or intramuscular injection, but oral or rectal administration is also contemplated. Ab and genetic vaccines are administered by injection, topically and orally. Dwg.0/16

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(FILE VOSPANEULLY ENTERED AT 12:52:32 ON 31 JUL 2002)
           1343 SEA FILE=HCAPLUS ABB=ON PLU=ON
                                                (MORAXEL? OR M OR
L1
                BRANHAMELL? OR M) (W) CATARRH?
             56 SEA FILE=HCAPLUS ABB=ON PLU=ON L1(5A)ANTIGEN
L4
             31 SEA FILE=HCAPLUS ABB=ON PLU=ON L4(S) VACCIN?
rs
             14 SEA FILE=USPATFULL ABB=ON PLU=ON L8(L)(POLYPEPTIDE OR
L16
                PEPTIDE OR PROTEIN OR POLYPROTEIN)
             14 SEA FILE-USPATFULL ABB=ON PLU=ON L16(L) (ANTIBOD? OR
L17
                T(W) (CELL OR LYMPHOCYT?))
L17 ANSWER 1 OF 14 USPATFULL
                        2002:140865 USPATFULL
ACCESSION NUMBER:
                        Vaccines comprising oil bodies
TITLE:
                        Deckers, Harm M., Alberta, CANADA
INVENTOR(S):
                        Rooijen, Gijs Van, Alberta, CANADA
                        Boothe, Joseph, Alberta, CANADA
                        Goll, Janis, Alberta, CANADA
                        Moloney, Maurice M., Alberta, CANADA
                        Schryvers, Anthony B., Alberta, CANADA
                        Alcantara, Joenel, Alberta, CANADA
                        Hutchins, Wendy A., Alberta, CANADA
                             NUMBER
                                        KIND
                                                  DATE
                        US 2002071846
                                                20020613
                                          A1
PATENT INFORMATION:
                        US 2001-880901
                                         A1
                                                20010615 (9)
APPLICATION INFO.:
                        Continuation-in-part of Ser. No. US 2000-577147,
RELATED APPLN. INFO.:
                        filed on 24 May 2000, PENDING
                        Continuation-in-part of Ser. No. US 1999-448600,
                        filed on 24 Nov 1999, PATENTED
                        Continuation-in-part of Ser. No. US 1998-84777,
                        filed on 27 May 1998, PATENTED
                               NUMBER
                                             DATE
                             ______
                        US 1998-75863P
                                           19980225 (60)
PRIORITY INFORMATION:
                                           19980225 (60)
                        US 1998-75864P
                                           19970528 (60)
                        US 1997-47779P
                                           19970527 (60)
                        US 1997-47753P
                        US 2000-212130P
                                           20000616 (60)
DOCUMENT TYPE:
                        Utility
                        APPLICATION
FILE SEGMENT:
                        BURNS DOANE SWECKER & MATHIS L L P, POST OFFICE
LEGAL REPRESENTATIVE:
                        BOX 1404, ALEXANDRIA, VA, 22313-1404
NUMBER OF CLAIMS:
                        27
EXEMPLARY CLAIM:
                        1
NUMBER OF DRAWINGS:
                        10 Drawing Page(s)
                        2348
LINE COUNT:
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       The present invention provides novel adjuvants which comprise oil
AB
       bodies. The invention also provides vaccine formulations
       comprising oil bodies and an antigen and methods for preparing the
       vaccines and the use of the vaccines to elicit an immune response.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       INCLM: 424/184.100
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308-4994 Searcher : Shears

INCLS: 424/757.000; 424/731.000; 424/750.000; 424/758.000;

INCL

424/755.000; 424/764.000; 424/768.000

424/184.100 NCL NCLM:

424/757.000; 424/731.000; 424/750.000; 424/758.000; NCLS:

424/755.000; 424/764.000; 424/768.000

L17 ANSWER 2 OF 14 USPATFULL

ACCESSION NUMBER: 2002:115794 USPATFULL

Multi-component vaccine to protect against TITLE:

disease caused by Haemophilus influenzae and

(9)

Moraxella catarrhalis

Loosmore, Sheena M., Aurora, CANADA INVENTOR(S):

Yang, Yan-Ping, Willowdale, CANADA Klein, Michel H., Willowdale, CANADA

Sasaki, Ken, Willowdale, CANADA Aventis Pasteur Limited, Toronto, CANADA PATENT ASSIGNEE(S):

(non-U.S. corporation)

NUMBER KIND DATE

US 6391313 B1 20020521 PATENT INFORMATION: US 1999-353617 19990715

APPLICATION INFO.: Utility DOCUMENT TYPE:

GRANTED FILE SEGMENT:

PRIMARY EXAMINER: Graser, Jennifer E. LEGAL REPRESENTATIVE: Sim & McBurney

22 NUMBER OF CLAIMS: 1 EXEMPLARY CLAIM:

28 Drawing Figure(s); 18 Drawing Page(s) NUMBER OF DRAWINGS:

1437 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A multi-valent immunogenic composition confers protection on an AB immunized host against infection caused by both Haemophilus influenzae and Moraxella catarrhalis. Such composition comprises at least four antigens comprising at least one antigen from Haemophilus influenzae, and at least one antigen from Moraxella catarrhalis. Three of the antigens are adhesins. High molecular weight (HMW) proteins and Haemophilus influenzae adhesin (Hia) proteins of non-typeable Haemophilus and a 200 kDa outer membrane protein of Moraxella catarrhalis comprise the adhesin components while the other antigen is a non-proteolytic analog of Hin47 protein. Each component does not impair the immunogenicity of the others. The multi-valent immunogenic composition may be combined with DTP component vaccines, which may also include non-virulent poliovirus and PRP-T, to provide a component vaccine without impairment of the immunogenic properties of the other antigens.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

INCLM: 424/203.100 INCL

INCLS: 424/256.100; 424/251.100; 424/234.100; 424/193.100;

424/203.100; 424/197.110; 530/350.000

NCL NCLM: 424/203.100

424/193.100; 424/197.110; 424/234.100; 424/251.100; NCLS:

424/256.100; 530/350.000

L17 ANSWER 3 OF 14 USPATFULL

2001:191256 USPATFULL ACCESSION NUMBER:

USPA1 and USPA2 antigens of Moraxella catarrhalis TITLE:

Hansen, Eric J., Plano, TX, United States INVENTOR(S):

> 308-4994 Shears Searcher :

Aebi, Christoph, Gasel, Switzerland

Cope, Leslie D., Mesquite, TX, United States Maciver, Isobel, Cottage Grove, WI, United States Fiske, Michael J., Rochester, NY, United States Fredenburg, Ross A., Rochester, NY, United States

PATENT ASSIGNEE(S):

Board of Regents, The University of Texas, Austin, TX, United States (U.S. corporation) American Cyanamid, Madison, NJ, United States

(U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.: US 6310190 B1 20011030 US 1999-336447 19990621 (9)

Continuation of Ser. No. WO 1997-US23930, filed

on 19 Dec 1997

NUMBER DATE

PRIORITY INFORMATION:

US 1996-33598P 19961220 (60)

DOCUMENT TYPE: FILE SEGMENT:

Utility GRANTED

PRIMARY EXAMINER: ASSISTANT EXAMINER:

Jones, W. Gary Soudaya, Jehanne Fulbright & Jaworski

LEGAL REPRESENTATIVE: NUMBER OF CLAIMS: EXEMPLARY CLAIM:

2

NUMBER OF DRAWINGS:

28 Drawing Figure(s); 17 Drawing Page(s)

LINE COUNT: 4794.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention discloses the existence of two novel proteins UspA1 and UspA2, and their respective genes uspA1 and uspA2. Each protein encompasses a region that is conserved between the two proteins and comprises an epitope that is recognized by the MAb 17C7. One or more than one of these species may aggregate to form the very high molecular weight form (i.e. greater than 200 kDa) of the UspA antigen. Compositions and both diagnostic and therapeutic methods for the treatment and study of M. catarrhalis are disclosed.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

INCL INCLM: 536/023.100 INCLS: 536/023.700

NCL NCLM: 536/023.100 NCLS: 536/023.700

PATENT ASSIGNEE(S):

L17 ANSWER 4 OF 14 USPATFULL

ACCESSION NUMBER: 2001:157808 USPATFULL

TITLE: Transferrin receptor protein of Moraxella

Tunistatin receptor process or notes

INVENTOR(S): Yang, Yan-Ping, Willowdale, Canada Myers, Lisa E., Guelph, Canada

Harkness, Robin E., Willowdale, Canada

Klein, Michel H., Willowdale, Canada Aventis Pasteur Limited, Toronto, Canada

(non-U.S. corporation)

. NUMBER KIND DATE

PATENT INFORMATION: US 6290970 B1 20010918 US 1995-540753 19951011 (8) APPLICATION INFO.: Utility DOCUMENT TYPE: GRANTED FILE SEGMENT: PRIMARY EXAMINER: Minnifield, Nita Sim & McBurney LEGAL REPRESENTATIVE: NUMBER OF CLAIMS: 7 EXEMPLARY CLAIM: 12 Drawing Figure(s); 8 Drawing Page(s) NUMBER OF DRAWINGS: LINE COUNT: 1199 CAS INDEXING IS AVAILABLE FOR THIS PATENT. An isolated and purified non-denatured transferrin receptor protein of a Moraxella strain, particularly M. catarrhalis, has an apparent molecular mass of about 80 to about 90 kDa, as determined by SDS-PAGE. The transferrin receptor protein or a fragment analog thereof is useful in diagnostic applications and immunogenic compositions, particularly for in vivo administration to a host to confer protection against disease caused by a strain of Moraxella. CAS INDEXING IS AVAILABLE FOR THIS PATENT. INCLM: 424/251.100 INCL INCLS: 530/350.000; 530/412.000; 424/190.100; 424/250.100; 424/184.100; 424/234.100; 514/002.000 NCL NCLM: 424/251.100 424/184.100; 424/190.100; 424/234.100; 424/250.100; NCLS: 514/002.000; 530/350.000; 530/412.000 L17 ANSWER 5 OF 14 USPATFULL ACCESSION NUMBER: 2001:52204 USPATFULL Moraxella catarrhalis outer membrane protein-106 TITLE: polypeptide, gene sequence and uses thereof INVENTOR(S): Tucker, Kenneth, Frederick, MD, United States Plosila, Laura, Cary, NC, United States Tillman, Ulrich F., Olney, MD, United States PATENT ASSIGNEE(S): Antex Biologics Inc., Gaithersburg, MD, United States (U.S. corporation) NUMBER · KIND DATE US 6214981 В1 20010410 PATENT INFORMATION: 19971112 APPLICATION INFO.: US 1997-968685 (8) Continuation-in-part of Ser. No. US 1996-642712, RELATED APPLN. INFO.: filed on 3 May 1996 DOCUMENT TYPE: Utility FILE SEGMENT: Granted PRIMARY EXAMINER: Smith, Lynette R. F. ASSISTANT EXAMINER: Portner, Ginny Allen Pennie & Edmonds LLP LEGAL REPRESENTATIVE: NUMBER OF CLAIMS: EXEMPLARY CLAIM: 15 Drawing Figure(s); 13 Drawing Page(s) NUMBER OF DRAWINGS: LINE COUNT: 2357 CAS INDEXING IS AVAILABLE FOR THIS PATENT. The invention discloses the Moraxella catarrhalis outer membrane protein-106 (OMP106) polypeptide, polypeptides derived therefrom (OMP106-derived polypeptides), nucleotide sequences encoding said polypeptides, and antibodies that specifically bind the OMP106

Searcher: Shears 308-4994

polypeptide and/or OMP106-derived polypeptides. Also disclosed are

immunogenic, prophylactic or therapeutic compositions, including vaccines, comprising OMP106 polypeptide and/or OMP106-derived polypeptides. The invention additionally discloses methods of inducing immune responses to M. catarrhalis and M. catarrhalis OMP106 polypeptides and OMP106-derived polypeptides in animals.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

INCL INCLM: 536/023.100

INCLS: 536/023.700; 424/184.100; 424/190.100; 424/234.100

NCL NCLM: 536/023.100

NCLS: 424/184.100; 424/190.100; 424/234.100; 536/023.700

L17 ANSWER 6 OF 14 USPATFULL

ACCESSION NUMBER: 2001:25435 USPATFULL

TITLE: Transferrin receptor protein of moraxella

INVENTOR(S): Yang, Yan-Ping, Willowdale, Canada

Myers, Lisa E., Guelph, Canada

Harkness Robin F. Willowdale, Canada

Harkness, Robin E., Willowdale, Canada Klein, Michel H., Willowdale, Canada

PATENT ASSIGNEE(S): Connaught Laboratories Limited, Toronto, Canada

(non-U.S. corporation)

KIND DATE NUMBER B1 20010220 PATENT INFORMATION: US 6190668 19970417 WO 9713785 19980730 (9)US 1998-51320 APPLICATION INFO .: 19961011 WO 1996-CA684 19980730 PCT 371 date 19980730 PCT 102(e) date

RELATED APPLN. INFO.: Continuation of Ser. No. US 1995-540753, filed on

11 Oct 1995

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Minnifield, Nita LEGAL REPRESENTATIVE: Sim & McBurney

NUMBER OF CLAIMS: 8
EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 5 Drawing Figure(s); 8 Drawing Page(s)

LINE COUNT: 1221

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

An isolated and purified non-denatured transferrin receptor protein of a Moraxella strain, particularly M. catarrhalis, has an apparent molecular mass of about 80 to about 90 kDa, as determined by SDS-PAGE. The transferrin receptor protein or a fragment analog thereof is useful in diagnostic applications and immunogenic compositions, particularly for in vivo administration to a host to confer protection against disease caused by a strain of Moraxella. The transferrin receptor protein is isolated from strains of Moraxella catarrhalis by a procedure including extraction of agent soluble proteins of a cell mass produced by cultivating the strain under iron-starved conditions. The transferrin receptor protein is selectively solubilized from the extracted cell mass and purified.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

INCL INCLM: 424/251.100

INCLS: 530/387.100; 530/412.000; 530/417.000; 435/007.100; 435/007.800; 435/070.200

NCL NCLM: 424/251.100

NCLS: 435/007.100; 435/007.800; 435/070.200; 530/387.100;

530/412.000; 530/417.000

L17 ANSWER 7 OF 14 USPATFULL

ACCESSION NUMBER: 2001:18617 USPATFULL

TITLE: Lactoferrin receptor genes of Moraxella

INVENTOR(S): Loosmore, Sheena M., Aurora, Canada

Du, Run-Pan, Thornhill, Canada Wang, Quijun, Thornhill, Canada Yang, Yan-Ping, Willowdale, Canada Klein, Michel H., Willowdale, Canada

PATENT ASSIGNEE(S): Connaught Laboratories Limited, Toronto, Canada

(non-U.S. corporation)

PATENT INFORMATION: US 6184371 B1 20010206 APPLICATION INFO.: US 1998-74658 19980508 (9)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1997-867941,

filed on 3 Jun 1997, now patented, Pat. No. US

5977337 Utility

DOCUMENT TYPE: Utility
FILE SEGMENT: Granted

PRIMARY EXAMINER: Graser, Jennifer LEGAL REPRESENTATIVE: Sim & McBurney

NUMBER OF CLAIMS: 5 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 140 Drawing Figure(s); 130 Drawing Page(s)

LINE COUNT: 1824

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Purified and isolated nucleic acid molecules are provided which encode lactoferrin receptor proteins of Moraxella, such as M. catarrhalis, or a fragment or an analog of the lactoferrin receptor protein. The nucleic acid sequence may be used to produce recombinant lactoferrin receptor proteins Lbp1, Lbp2 and ORF3 of the strain of Moraxella free of other proteins of the Moraxella strain for purposes of diagnostics and medical treatment. Furthermore, the nucleic acid molecule may be used in the diagnosis of infection.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

INCL INCLM: 536/023.700

INCLS: 536/023.100; 536/024.300; 536/024.320; 435/320.100; 435/069.100; 435/069.300; 435/069.700; 435/252.300;

424/200.100; 424/251.100

NCL NCLM: 536/023.700

NCLS: 424/200.100; 424/251.100; 435/069.100; 435/069.300; 435/069.700; 435/252.300; 435/320.100; 536/023.100;

536/024.300; 536/024.320

L17 ANSWER 8 OF 14 USPATFULL

ACCESSION NUMBER: 1999:166603 USPATFULL

TITLE: Outer membrane protein B1 of Moraxella

catarrhalis

INVENTOR(S): Campagnari, Anthony A., Hamburg, NY, United

States

PATENT ASSIGNEE(S): The Research Foundation of the State University

of New York, Amherst, NY, United States (U.S. corporation)

	NUMBER	KIND	DATE	
•				
PATENT INFORMATION:	US 6004562		19991221	
APPLICATION INFO .:	US 1996-698652		19960816	(8)
DOCUMENT TYPE.	FT4-2-3-2-4			

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Housel, James C.

ASSISTANT EXAMINER: Ryan, V.

Hodgson, Russ, Andrews, Woods & Goodyear, LLP LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: 10 EXEMPLARY CLAIM: 1

3 Drawing Figure(s); 2 Drawing Page(s) NUMBER OF DRAWINGS:

LINE COUNT: 915

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

An isolated and purified outer membrane protein B1, and peptides formed therefrom, of Moraxella catarrhalis are described. A method for the isolation and purification of outer membrane protein B1 from a bacterial strain that produces B1 protein, e.g. Moraxella catarrhalis, comprises growing the bacteria in culture in iron-depleted medium to enhance the expression of the B1 protein, harvesting the bacteria from the culture, extracting from the harvested bacteria a preparation substantially comprising an outer membrane protein preparation, contacting the outer membrane preparation with an affinity matrix containing immobilized transferrin wherein B1 protein binds to the transferrin, and eluting the bound B1 protein from the transferrin. Disclosed are the uses of the B1 protein as an immunogen for vaccine formulations, and as antigens in diagnostic immunoassays.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

INCL INCLM: 424/251.100

INCLS: 424/184.100; 424/234.100

NCL NCLM: 424/251.100

DOCUMENT TYPE:

424/184.100; 424/234.100 NCLS:

L17 ANSWER 9 OF 14 USPATFULL

1999:155210 USPATFULL ACCESSION NUMBER:

Methods and compositions relating to useful TITLE:

antigens of moraxella catarrhalis

Hansen, Eric J., Plano, TX, United States INVENTOR(S):

Helminen, Meria E., Helsinki, Finland

Maciver, Isobel, Dallas, TX, United States Board of Regents, The University of Texas,

PATENT ASSIGNEE(S): Austin, TX, United States (U.S. corporation)

	NUMBER	KIND	DATE	
PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:	US 5993826 US 1993-25363 Continuation-in- filed on 14 Aug continuation-in- filed on 21 Aug 5552146	part of 1992 whi part of	ch is a Ser. No.	wo 1992-US6869, US 1991-745591,

Searcher : Shears 308-4994

Utility

FILE SEGMENT:

Granted

PRIMARY EXAMINER: LEGAL REPRESENTATIVE: Sidberry, Hazel F. Arnold White & Durkee

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 11

NUMBER OF DRAWINGS:

19 Drawing Figure(s); 17 Drawing Page(s)

LINE COUNT: 3037

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB

The present disclosure relates to Moraxella catarrhalis outer membrane vesicle (OMV) compositions, to selected antigenic proteins from the outer membranes of M. catarrhalis which have a variety of useful properties, and to monoclonal antibodies against these proteins. Particular "Outer Membrane Proteins" (OMPs) of the invention are characterized as having molecular weights of about 30 kD, 80 kD (also termed CopB protein) and between about 200 and 700 kD (HMWP antigen). Passive immunization with monoclonal antibodies directed against these proteins confers protection against homologous and heterologous Moraxella catarrhalis strains in animal models, and active immunization with outer membrane vesicles also enhances pulmonary clearance of distinct M. catarrhalis strains. This demonstrates both the utility of antibodies in conferring passive immunity and the usefulness of OMPs, or variants thereof, in the preparation of vaccines. Also disclosed are DNA segments encoding these OMPs, methods for preparing the antigens, or variants, through the application of recombinant DNA techniques, as well as diagnostic methods and related embodiments.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

INCL INCLM: 424/251.100

INCLS: 424/184.100; 530/350.000; 530/388.100; 530/388.200;

435/069.100; 435/069.300

NCL NO

NCLM: 424/251.100

NCLS: 424/184.100; 435/069.100; 435/069.300; 530/350.000;

530/388.100; 530/388.200

L17 ANSWER 10 OF 14 USPATFULL

ACCESSION NUMBER:

1999:141620 USPATFULL

TITLE:

Methods and compositions relating to useful

antigens of moraxella catarrhalis

INVENTOR(S):

Hansen, Eric J., Plano, TX, United States Helminen, Merja E., Helsinki, Finland Maciver, Isobel, Dallas, TX, United States

PATENT ASSIGNEE(S):

Board of Regents, The University of Texas System,

Austin, TX, United States (U.S. corporation)

	NUMBER	KIND	DATE
,			
PATENT INFORMATION:	US 5981213		19991109
ADDITONTION THEO .	rrs 1995-450351		19950525

APPLICATION INFO.: RELATED APPLN. INFO.:

Division of Ser. No. US 1993-25363, filed on 2 Mar 1993 which is a continuation-in-part of Ser. No. WO 1992-US6869, filed on 14 Aug 1992, now patented, Pat. No. WO 819315, issued on 19 Sep 1994 which is a continuation-in-part of Ser. No.

US 1991-745591, filed on 21 Aug 1991, now

patented, Pat. No. US 5552146

DOCUMENT TYPE:

Utility

FILE SEGMENT:

Granted

PRIMARY EXAMINER: ASSISTANT EXAMINER: Housel, James C. Shaver, Jennifer

LEGAL REPRESENTATIVE:

Arnold, White & Durkee

NUMBER OF CLAIMS:

23

EXEMPLARY CLAIM:

1

NUMBER OF DRAWINGS:

13 Drawing Figure(s); 17 Drawing Page(s)

LINE COUNT:

3099

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present disclosure relates to Moraxella catarrhalis outer membrane vesicle (OMV) compositions, to selected antigenic proteins from the outer membranes of M. catarrhalis which have a variety of useful properties, and to monoclonal antibodies against these proteins. Particular "Outer Membrane Proteins" (OMPs) of the invention are characterized as having molecular weights of about 30 kD, 80 kD (also termed CopB protein) and between about 200 and 700 kD (HMWP antigen). Passive immunization with monoclonal antibodies directed against these proteins confers protection against homologous and heterologous Moraxella catarrhalis strains in animal models, and active immunization with outer membrane vesicles also enhances pulmonary clearance of distinct M. catarrhalis strains. This demonstrates both the utility of antibodies in conferring passive immunity and the usefulness of OMPs, or variants thereof, in the preparation of vaccines. Also disclosed are DNA segments encoding these OMPs, methods for preparing the antigens, or variants, through the application of recombinant DNA techniques, as well as diagnostic methods and related embodiments.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

INCL

INCLM: 435/069.100

INCLS: 435/069.300; 435/252.200; 435/320.100; 536/023.100; 536/023.700; 536/024.320; 424/234.100; 424/251.100

NCL

NCLM: 435/069.100

424/234.100; 424/251.100; 435/069.300; 435/252.200;

435/320.100; 536/023.100; 536/023.700; 536/024.320

L17 ANSWER 11 OF 14 USPATFULL

ACCESSION NUMBER:

1999:106092 USPATFULL

TITLE:

Vaccine for Moraxella catarrhalis

INVENTOR(S):

Murphy, Timothy F., East Amherst, NY, United

States

PATENT ASSIGNEE(S):

The Research Foundation of State University of New York, Amherst, NY, United States (U.S.

corporation)

NUMBER KIND DATE 19990907

PATENT INFORMATION:

US 5948412 US 1997-810655

APPLICATION INFO.: RELATED APPLN. INFO.:

Continuation-in-part of Ser. No. US 1994-245758, filed on 17 May 1994, now patented, Pat. No. US

19970303

5607846

DOCUMENT TYPE: FILE SEGMENT:

Utility Granted

PRIMARY EXAMINER:

Degen, Nancy

ASSISTANT EXAMINER: LEGAL REPRESENTATIVE: Schwartzman, Robert

Hodgson, Russ, Andrews Woods & Goodyear, LLP

Searcher :

Shears

308-4994

(8)

NUMBER OF CLAIMS: 17 EXEMPLARY CLAIM: 1

3 Drawing Figure(s); 2 Drawing Page(s) NUMBER OF DRAWINGS:

LINE COUNT: 1552

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Compositions comprising outer membrane protein "E", and peptides AB and oligopeptides thereof, of Moraxella catarrhalis are described. Additionally, nucleotide sequences encoding the protein, peptide, or oligopeptide are disclosed, as well as recombinant vectors containing these sequences. Protein, peptide, or oligopeptide can be produced from host cell systems containing these recombinant vectors. Peptides and oligopeptides can also be chemically synthesized. Disclosed are the uses of the protein, peptides and oligopeptides as antigens in antigenic formulations for vaccine applications or for generating antisera of diagnostic or therapeutic use; and as antigens in diagnostic immunoassays. The nucleotide sequences are useful for constructing vectors for use as vaccines for insertions into attenuated bacteria in constructing a recombinant bacterial vaccine and for inserting into a viral vector in constructing a recombinant viral vaccine. Also described is the use of nucleotide sequences related to the gene encoding E as primers and/or probes in molecular diagnostic assays for the detection of M. catarrhalis.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

INCLM: 424/251.100 INCL INCLS: 530/350.000 NCLM: 424/251.100 NCL NCLS: 530/350.000

L17 ANSWER 12 OF 14 USPATFULL

ACCESSION NUMBER: 1998:61433 USPATFULL

Methods and compositions relating to useful TITLE:

antigens of moraxella catarrhalis

INVENTOR(S):

Hansen, Eric J., Plano, TX, United States Maciver, Isobel, Dallas, TX, United States

Helminen, Merja, Helsinki, Finland

Board of Regents, The University of Texas System, PATENT ASSIGNEE(S):

United States (U.S. corporation)

NUMBER KIND DATE 19980602 US 5759813 PATENT INFORMATION: US 1994-193150 APPLICATION INFO.: 19940919 (8)

Continuation of Ser. No. US 1991-745591, filed on RELATED APPLN. INFO.:

15 Aug 1991, now patented, Pat. No. US 5552146

DOCUMENT TYPE: Utility

Granted FILE SEGMENT:

Hutzell, Paula K. PRIMARY EXAMINER: Navarro, Mark ASSISTANT EXAMINER:

Arnold, White & Durkee LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: 15 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 5 Drawing Figure(s); 3 Drawing Page(s)

LINE COUNT: 1732

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present disclosure relates to selected antigenic proteins AB obtained from the outer membranes of Moraxella catarrhalis, that

are found to have a variety of useful properties. These proteins, termed OMPs ("Outer Membrane Proteins"), are characterized as having molecular weights of about 30 kD, 80 kD and between about 200 and 700 kD, respectively. Studies set forth herein demonstrated that monoclonal antibodies directed against these proteins confer a protective effect against infection by Moraxella catarrhalis organisms in animal models, demonstrating the potential usefulness of such antibodies in conferring passive immunity as well as the potential usefulness of these OMPs, or variants thereof, in the preparation of vaccines. Also disclosed are DNA segments encoding these OMPs, methods for preparing the antigens or variants, through the application of recombinant DNA techniques, as well as diagnostic methods and embodiments.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

INCL INCLM: 435/069.300

INCLS: 435/069.100; 435/320.100; 435/325.000; 536/023.100;

536/023.700; 530/350.000; 424/184.100

NCL NCLM: 435/069.300

NCLS: 424/184.100; 435/069.100; 435/320.100; 435/325.000;

530/350.000; 536/023.100; 536/023.700

L17 ANSWER 13 OF 14 USPATFULL

ACCESSION NUMBER: 97:9925 USPATFULL

TITLE: Methods and compositions relating to useful

antigens of moraxella catarrhalis

INVENTOR(S): Hansen, Eric J., Plano, TX, United States

Helminen, Merja, Dallas, TX, United States Maciver, Isobel, Dallas, TX, United States

PATENT ASSIGNEE(S): American Cyanamid Company, Wayne, NJ, United

States (U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 5599693 19970204 APPLICATION INFO.: US 1995-450002 19950525 (8)

RELATED APPLN. INFO.: Division of Ser. No. US 1991-745591, filed on 15

Aug 1991

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Housel, James C. ASSISTANT EXAMINER: Murthy, Prasad

LEGAL REPRESENTATIVE: Arnold White & Durkee

NUMBER OF CLAIMS: 12 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 3 Drawing Figure(s); 2 Drawing Page(s)

LINE COUNT: 1620

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present disclosure relates to selected antigenic proteins obtained from the outer membranes of Moraxella catarrhalis, that have been found by the inventors to have a variety of useful properties. These proteins, termed OMPs ("Outer Membrane Proteins"), are characterized as having molecular weights of 30, 80 and 100 kD, respectively. Studies set forth herein demonstrate that monoclonal antibodies directed against these proteins confer a protective effect against infection by Moraxella catarrhalis organisms in animal models, demonstrating the potential usefulness of such antibodies in conferring passive immunity as well as the

potential usefulness of these OMPs, or variants thereof, in the preparation of vaccines. Also disclosed are DNA segments encoding these OMPs, methods for preparing the antigens, or variants, through the application of recombinant DNA techniques, as well as diagnostic methods and embodiments.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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INCL
       INCLM: 435/069.300
       INCLS: 424/184.100; 424/251.100; 435/007.200; 435/007.320;
              435/071.100; 435/071.200; 435/243.000; 435/252.100;
              436/543.000; 530/388.200; 530/388.400; 530/412.000;
              530/413.000; 935/106.000; 935/108.000; 935/109.000;
              935/110.000
NCL
       NCLM:
              435/069.300
              424/184.100; 424/251.100; 435/007.200; 435/007.320;
      NCLS:
              435/071.100; 435/071.200; 435/243.000; 435/252.100;
              436/543.000; 530/388.200; 530/388.400; 530/412.000;
              530/413.000
L17 ANSWER 14 OF 14 USPATFULL
                         96:80017 USPATFULL
ACCESSION NUMBER:
                        Methods and compositions relating to useful
TITLE:
                         antigens of Moraxella catarrhalis
INVENTOR(S):
                        Hansen, Eric J., Plano, TX, United States
                        Helminen, Merja, Dallas, TX, United States
Maciver, Isobel, Dallas, TX, United States
                        Board of Regents, The University of Texas System,
PATENT ASSIGNEE(S):
                        Austin, TX, United States (U.S. corporation)
                                           KIND
                                                   DATE
                              NUMBER
                         US 5552146
                                                 19960903
PATENT INFORMATION:
                         US 1991-745591
                                                 19910815
                                                            (7)
APPLICATION INFO.:
DOCUMENT TYPE:
                        Utility
FILE SEGMENT:
                        Granted
                        Sidberry, Hazel F.
PRIMARY EXAMINER:
                        Arnold, White & Durkee
LEGAL REPRESENTATIVE:
NUMBER OF CLAIMS:
EXEMPLARY CLAIM:
                        3 Drawing Figure(s); 2 Drawing Page(s)
NUMBER OF DRAWINGS:
                        1597
LINE COUNT:
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       The present disclosure relates to selected antigenic proteins
       obtained from the outer membranes of Moraxella catarrhalis, that
       have been found by the inventors to have a variety of useful
       properties. These proteins, termed OMPs ("Outer Membrane
       Proteins"), are characterized as having molecular weights of 30,
       80 and 100 kD, respectively. Studies set forth herein demonstrate
       that monoclonal antibodies directed against these proteins confer
       a protective effect against infection by Moraxella catarrhalis
       organisms in animal models, demonstrating the potential usefulness
       of such antibodies in conferring passive immunity as well as the
       potential usefulness of these OMPs, or variants thereof, in the
       preparation of vaccines. Also disclosed are DNA segments encoding
       these OMPs, methods for preparing the antigens, or variants,
       through the application of recombinant DNA techniques, as well as
       diagnostic methods and embodiments.
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CAS INDEXING IS AVAILABLE FOR THIS PATENT.

INCLM: 424/251.100

INCLS: 424/184.100; 530/350.000

424/251.100 NCL NCLM:

424/184.100; 530/350.000 NCLS:

(FILE MEDLINE ENTERED AT 12:55:00 ON 31 JUL 2002)

1021 SEA FILE=MEDLINE ABB=ON PLU=ON "MORAXELLA (BRANHAMELLA) L18

CATARRHALIS"/CT

VACCINES/CT 5674 SEA FILE=MEDLINE ABB=ON PLU=ON L19

29132 SEA FILE=MEDLINE ABB=ON PLU=ON VACCINATION/CT

9 SEA FILE=MEDLINE ABB=ON PLU=ON L18 AND (L19 OR L20)

PLU=ON "MORAXELLA (BRANHAMELLA) 1021 SEA FILE=MEDLINE ABB=ON L18

CATARRHALIS"/CT

47913 SEA FILE=MEDLINE ABB=ON PLU=ON ANTIGENS/CT

1 SEA FILE=MEDLINE ABB=ON PLU=ON L18 AND L22



L20

10-L21-OR-L23

L24 ANSWER 1 OF 10 MEDLINE

AN 2000428046 MEDLINE

Enhancement of clearance of bacteria from murine lungs by TI immunization with detoxified lipooligosaccharide from Moraxella. catarrhalis conjugated to proteins.

·Hu W G; Chen J; Battey J F; Gu X X ΑU

INFECTION AND IMMUNITY, (2000 Sep) 68 (9) 4980-5. Journal code: 0246127. ISSN: 0019-9567. SO

Moraxella catarrhalis strain 25238 detoxified lipooligosaccharide AΒ (dLOS)-protein conjugates induced a significant rise of bactericidal anti-LOS antibodies in animals. This study reports the effect of active or passive immunization with the conjugates or their antiserum on pulmonary clearance of M. catarrhalis in an aerosol challenge mouse model. Mice were injected subcutaneously with dLOS-tetanus toxoid (dLOS-TT), dLOS-high-molecular-weight proteins (dLOS-HMP) from nontypeable Haemophilus influenzae (NTHi), or nonconjugated materials in Ribi adjuvant and then challenged with M. catarrhalis strain 25238 or O35E or NTHi strain 12. Immunization with dLOS-TT or dLOS-HMP generated a significant rise of serum anti-LOS immunoglobulin G and 68% and 35 to 41% reductions of bacteria in lungs compared with the control (P<0.01) following challenge with homologous strain 25238 and heterologous strain 035E, respectively. Serum anti-LOS antibody levels correlated with its bactericidal titers against M. catarrhalis and bacterial CFU in lungs. Additionally, immunization with dLOS-HMP generated a 54% reduction of NTHi strain 12 compared with the control (P<0.01). Passive immunization with a rabbit antiserum against dLOS-TT conferred a significant reduction of strain 25238 CFU in lungs in a dose- and time-dependent pattern compared with preimmune serum-treated mice. Kinetic examination of lung tissue sections demonstrated that antiserum-treated mice initiated and offset inflammatory responses more rapidly than preimmune serum-treated mice. These data indicate that LOS antibodies (whether active or passive) play a major role in the enhancement of pulmonary clearance of different test strains of M. catarrhalis in mice. In addition, dLOS-HMP is a potential candidate for a bivalent vaccine against M.

catarrhalis and NTHi infections.

- L24 ANSWER 2 OF 10 MEDLINE
- AN 2000398416 MEDLINE
- TI Potential of bacterial vaccines in the prevention of acute otitis media.
- AU Eskola J; Kilpi T
- SO PEDIATRIC INFECTIOUS DISEASE JOURNAL, (2000 May) 19 (5 Suppl) S72-8. Ref: 83
 Journal code: 8701858. ISSN: 0891-3668.
- L24 ANSWER 3 OF 10 MEDLINE
- AN 1999458176 MEDLINE
- TI The promise of immunoprophylaxis for prevention of acute otitis media.
- AU Pelton S I; Klein J O
- SO PEDIATRIC INFECTIOUS DISEASE JOURNAL, (1999 Oct) 18 (10) 926-35. Ref: 92
 Journal code: 8701858. ISSN: 0891-3668.
- L24 ANSWER 4 OF 10 MEDLINE
- AN 1999000946 MEDLINE
- TI Otitis media: focus on antimicrobial resistance and new treatment options.
- AU Hoppe H L; Johnson C E
- SO AMERICAN JOURNAL OF HEALTH-SYSTEM PHARMACY, (1998 Sep 15) 55 (18) 1881-97; quiz 1932-3. Ref: 99 Journal code: 9503023. ISSN: 1079-2082.
- Antimicrobial resistance among organisms that cause acute otitis AB media (AOM) and new approaches in the prevention and treatment of AOM are discussed. Organisms commonly responsible for causing AOM include Streptococcus pneumoniae, Haemophilus influenzae, and Moraxella catarrhalis. The evolution of pneumococcal resistance to penicillins, erythromycin, trimethoprim-sulfamethoxazole, and oral cephalosporins may require treatment with agents such as vancomycin or rifampin in certain patients. H. influenzae and M. catarrhalis are becoming increasingly resistant to penicillins, trimethoprim-sulfamethoxazole, oral cephalosporins, and macrolides. Mechanisms of resistance include changes in penicillin-binding proteins, production of beta-lactamase, alterations in target enzymes, and inhibition of drug access to the site of action. Because of changing resistance patterns and the limited spectra of activity of many currently available antimicrobials, new antimicrobials have been developed in the hope of improving therapy. While amoxicillin and trimethoprim-sulfamethoxazole are appropriate first-line agents, children at risk for resistant infections may be treated initially with cefuroxime axetil, cefpodoxime proxetil, cefprozil, or amoxicillin-clavulanate. After local resistance patterns, patient adherence to therapy, in vitro data, and cost factors have been weighed, other agents to consider include loracarbef, clarithromycin, azithromycin, and ceftriaxone. Along with the efforts to improve treatment, research is focusing on the prevention of otitis media with bacterial and viral vaccines. The emergence of resistant strains of organisms causing AOM has complicated its treatment.
- L24 ANSWER 5 OF 10 MEDLINE
- AN 1998279666 MEDLINE

- TI Vaccination against middle-ear bacterial and viral pathogens.
- AU Giebink G S
- SO ANNALS OF THE NEW YORK ACADEMY OF SCIENCES, (1997 Dec 29) 830 330-52. Ref: 121 Journal code: 7506858. ISSN: 0077-8923.
- Considerable evidence suggests that otitis media (OM) can be AB prevented by systemic immunization. Building on the highly effective H. influenzae type b (Hib) conjugate vaccine technology, pneumococcal conjugate vaccines are being developed to circumvent T-independence of these antigens and provide durable immunity at a very young age. Several pneumococcal conjugate vaccines are currently in clinical testing. Potential vaccine antigens of nontypable H. influenzae (NTHi) include OMP, HMW, pili, and fimbriae. Several OMPs show extensive homology among strains, but surface, determinants of others are highly variable so that antibodies to surface epitopes of one strain will not bind to surface epitopes of another. Several M. catarrhalis OMP and HMW antigens have vaccine potential, but no functional correlates of protection have been identified, and there is no clear evidence that antibody to M. catarrhalis is associated with OM protection. Attenuated viral vaccines also hold promise of preventing childhood OM. Two clinical trials with killed influenza vaccines have shown a significant reduction in OM among vaccine recipients compared to control children during periods of high influenza disease activity in the community. Passive immunoprophylaxis also has potential for preventing OM. Human bacterial polysaccharide immune globulin was protective for pneumococcal OM in children and in the chinchilla OM model. High-dose respiratory syncytial virus-enriched immunoglobulin reduced the incidence and severity of RSV lower respiratory tract infection in high-risk children. Passive immunoprophylaxis may also be effective in children with specific immune deficiencies, such as IgG2 deficiency, and patients who fail to respond to vaccines.
- L24 ANSWER 6 OF 10 MEDLINE
- AN 97130436 MEDLINE
- TI Dendritic cells are recruited into the airway epithelium during the inflammatory response to a broad spectrum of stimuli.
- AU McWilliam A S; Napoli S; Marsh A M; Pemper F L; Nelson D J; Pimm C L; Stumbles P A; Wells T N; Holt P G
- SO JOURNAL OF EXPERIMENTAL MEDICINE, (1996 Dec 1) 184 (6) 2429-32. Journal code: 2985109R. ISSN: 0022-1007.
- A key rate-limiting step in the adaptive immune response at AB peripheral challenge sites is the transmission of antigen signals to T cells in regional lymph nodes. Recent evidence suggests that specialized dendritic cells (DC) fulfill this surveillance function in the resting state, but their relatively slow turnover in most peripheral tissues brings into question their effectiveness in signaling the arrival of highly pathogenic sources of antigen which require immediate mobilization of the full range of host defenses for maintenance of homeostasis. However, the present report demonstrates that recruitment of a wave of DC into the respiratory tract mucosa is a universal feature of the acute cellular response to local challenge with bacterial, viral, and soluble protein antigens. Consistent with this finding, we also demonstrate that freshly isolated respiratory mucosal DC respond in vitro to a variety of CC chemokines as well as complementary cleavage products and N-formyl-methionyl-leucine-phenylalanine. This suggests that rapid amplification of specific antigen surveillance at peripheral

challenge sites is an integral feature of the innate immune response at mucosal surfaces, and serves as an "early warning system" to alert the adaptive immune system to incoming pathogens.

- L24 ANSWER 7 OF 10 MEDLINE
- AN 96238995 MEDLINE
- TI Evaluation of purified UspA from Moraxella catarrhalis as a vaccine in a murine model after active immunization.
- AU Chen D; McMichael J C; VanDerMeid K R; Hahn D; Mininni T; Cowell J; Eldridge J
- SO INFECTION AND IMMUNITY, (1996 Jun) 64 (6) 1900-5. Journal code: 0246127. ISSN: 0019-9567.
- Moraxella catarrhalis causes otitis media, laryngitis, and AB: respiratory infections in humans. A high-molecular-weight outer membrane protein from this bacterium named ubiquitous surface protein A (UspA) is present on all isolates. A monoclonal antibody (MAb) to UspA that recognizes a conserved epitope of this protein has been shown to promote pulmonary clearance of bacteria in passively immunized mice. In the present study, M. catarrhalis heterologous isolates were screened by dot blot with a panel of four additional MAbs specific for surface-exposed epitopes of UspA from M. catarrhalis isolate 035E. Three of the MAbs were specific for 035E, and the fourth reacted with 17 (74%) of the 23 isolates tested. Thus, UspA contains highly conserved, semiconserved, and variable surface-exposed epitopes. The UspA was purified from the 035E isolate by ion-exchange and size-exclusion chromatography, formulated with the adjuvant QS-21, and used to immunize BALB/c mice. Upon pulmonary challenge with either 035E or the heterologous isolate TTA24, significantly fewer bacteria were recovered from the lungs of immunized mice 6 h postchallenge than from control mice. The immune sera from mice or guinea pigs contained high titers of antibodies to the homologous isolate and heterologous isolates in a whole-bacterial-cell enzyme-linked immunosorbent assay. Sera against UspA, whether prepared in mice or guinea pigs, had complement-dependent bactericidal activity toward homologous and 11 heterologous M. catarrhalis isolates. These results indicate that the conserved epitopes of the UspA are highly immunogenic and elicit broadly reactive and biologically functional antibodies. UspA may offer protection against M. catarrhalis infections and is being further evaluated as a vaccine candidate.
- L24 ANSWER 8 OF 10 MEDLINE
- AN 94234646 MEDLINE
- TI Preventing otitis media.
- AU Giebink G S
- SO ANNALS OF OTOLOGY, RHINOLOGY, AND LARYNGOLOGY. SUPPLEMENT, (1994 May) 163 20-3. Ref: 17
 Journal code: 1256156. ISSN: 0096-8056.
- AB Recurrent acute otitis media (AOM) is an extremely prevalent disease in young children. Epidemiologic associations suggest that primary prevention or reduction of AOM frequency may be achieved with breast-feeding during infancy, elimination of household tobacco smoking, and use of small rather than large day-care arrangements for infants and toddlers. Secondary antimicrobial prophylaxis with amoxicillin or sulfisoxazole reduces the frequency of recurrent AOM by about 50%, but it does not appear to reduce the duration of otitis media with effusion (OME). Tympanostomy tube insertion is not as effective as amoxicillin in reducing AOM frequency in children

without OME. Adenoidectomy appears to be warranted for children who develop recurrent AOM after extrusion of tubes. Vaccines against the common bacteria and viruses causing AOM hold the greatest promise of preventing AOM and blocking the sequence of pathologic events leading to chronic OME and middle ear sequelae. The greatest progress has been made recently with pneumococcal protein conjugate vaccines, and clinical testing is in progress.

- L24 ANSWER 9 OF 10 MEDLINE
- AN 93329207 MEDLINE
- TI Effect of immunization of pulmonary clearance of Moraxella catarrhalis in an animal model.
- AU Maciver I; Unhanand M; McCracken G H Jr; Hansen E J
- SO JOURNAL OF INFECTIOUS DISEASES, (1993 Aug) 168 (2) 469-72. Journal code: 0413675. ISSN: 0022-1899.
- A murine model for pulmonary clearance of Moraxella catarrhalis was AB used to determine whether immunization could enhance clearance of this organism from the lungs. Animals actively immunized with outer membrane vesicles of M. catarrhalis cleared an endobronchial challenge with the homologous strain more quickly than did sham-immunized control animals. Western blot analysis of both this immune mouse serum and rabbit antiserum raised against outer membrane vesicles of M. catarrhalis indicated that antibodies were present to both outer membrane protein and lipooligosaccharide antigens. Passive immunization of mice with the immune rabbit serum resulted in enhanced pulmonary clearance of both homologous and heterologous strains of M. catarrhalis, indicating the involvement of serum antibody in this clearance process and the existence of conserved surface antigens in the two different M. catarrhalis strains. These results suggest that this model system may be useful for the identification of vaccine candidates among the surface antigens of M. catarrhalis.
- L24 ANSWER 10 OF 10 MEDLINE
- AN 93235586 MEDLINE
- TI Secretory IgA-, IgG- and C3b-coated bacteria in the nasopharynx of otitis-prone and non-otitis-prone children.
- AU Stenfors L E; Raisanen S
- SO ACTA OTO-LARYNGOLOGICA, (1993 Mar) 113 (2) 191-5. Journal code: 0370354. ISSN: 0001-6489.
- AB The proportions of secretory IgA (SIgA)-, IgG- and C3b-coated bacteria obtained from a well-defined area on the posterior wall of the nasopharynx (NPH) close to the Eustachian tube were determined. Samples taken from 25 otitis-prone (OP) and 25 non-otitis-prone (NOP) children with normal serum levels of IgA and IgG were evaluated using an immunofluorescence assay. Both groups harboured significantly more nasopharyngeal bacteria coated with IgG than with SIGA (p < 0.001). The OP children had significantly fewer SIgA-coated bacteria (p < 0.05) but more C3b-coated bacteria (p < 0.01) in the NPH than the NOP children had. No significant difference was noted between the two groups regarding IgG coating. The occurrence of Branhamella catarrhalis in the NHP was more pronounced in the OP group (p < 0.05). No significant differences in the occurrence of other middle ear pathogens (Streptococcus pneumoniae, Haemophilus influenzae, Staphylococcus aureus) or quantitative dominance of pathogens were noted between the two groups. Deficiency in SIgA coating of the nasopharyngeal bacteria may contribute to the otitis-prone condition.

(FLLE MCAPLUS, MEDLINE, BIOSIS, EMBASE, WPIDS, CONFSCI, SCISEARCH, JICST-EPLUS, JAPIO, USPATFULL' ENTERED AT 12:58:06 ON 31 JUL 2002)

22 S THONNARD J?/AU AND L8

20 DOP REM L25 (2 DUPLICATES REMOVED)

L26 ANSWER 1 OF 20 HCAPLUS COPYRIGHT 2002 ACS DUPLICATE 1

ACCESSION NUMBER: 2001:101183 HCAPLUS DOCUMENT NUMBER: 134:161878

TITLE: Moraxella catarrhalis BASB114 antigens and uses

thereof

INVENTOR(S):
Thonnard, Joelle

PATENT ASSIGNEE(S): Smithkline Beecham Biologicals S.A., Belg.

SOURCE: PCT Int. Appl., 82 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

DATE APPLICATION NO. PATENT NO. KIND DATE 20010208 WO 2000-EP7293 20000727 WO 2001009179 A1 AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG BF, BJ, EP 2000-956338 20000727 20020515 EP 1204678 A1 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, MC, IE, SI, LT, LV, FI, RO, MK, CY, AL GB 1999-17977 A 19990730 PRIORITY APPLN. INFO.: WO 2000-EP7293 W 20000727

AB The invention provides BASB114 polypeptides and polynucleotides encoding BASB114 polypeptides and methods for producing such polypeptides by recombinant techniques. Also provided are

diagnostic, prophylactic and therapeutic uses.

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN

THE RE FORMAT

L26 ANSWER 2 OF 20 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 2001:618174 HCAPLUS

DOCUMENT NUMBER: 135:191336

INVENTOR(S):

TITLE: Recombinant Haemophilus influenza outer membrane

protein and use thereof in vaccination Berthet, Francois-Xavier Jacques; Denoel, Philippe; Poolman, Jan; Thonnard, Joelle

PATENT ASSIGNEE(S): SmithKline Beecham Biologicals S.A., Belg.

SOURCE: PCT Int. Appl., 29 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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KIND DATE
                                             APPLICATION NO.
                                                               DATE
     PATENT NO.
                                             -----
                       ____
                             _____
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                                         WO 2001-EP1556 20010213
     WO 2001061013
                      A1 20010823
       W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH,
            CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ,
             PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU,
             TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH,
             CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
             TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD,
             TG
                                          GB 2000-3502
                                                            A 20000215
PRIORITY APPLN. INFO.:
     This invention relates to recombinant bacterial outer membrane
     proteins comprising one or more LB1(f) peptides from surface-exposed
     loop 3 of MOMP P5 of non-typeable H. influenzae. The invention also
     relates to a method of isolating the recombinant proteins and a
     vaccine compn. for use in the treatment of Haemophilus influenzae
     infection.
                                 THERE ARE 3 CITED REFERENCES AVAILABLE FOR
                          3
REFERENCE COUNT:
                                 THIS RECORD. ALL CITATIONS AVAILABLE IN
                                 THE RE FORMAT
                     WPIDS (C) 2002 THOMSON DERWENT
L26 ANSWER 3 OF 20
                       2001-244783 [25]
                                          WPIDS
ACCESSION NUMBER:
                       N2001-174285
DOC. NO. NON-CPI:
                       C2001-073454
DOC. NO. CPI:
                       Novel BASB129-BASB131 polypeptides isolated from
TITLE:
                       Moraxella catarrhalis bacterium useful as a
                       diagnostic reagent for M. catarrhalis infections and
                       for producing vaccines against otitis media and
                       pneumonia.
                       B04 D16 S03
DERWENT CLASS:
                       THONNARD, J
INVENTOR(S):
                       (SMIK) SMITHKLINE BEECHAM BIOLOGICALS
PATENT ASSIGNEE(S):
                       95
COUNTRY COUNT:
PATENT INFORMATION:
                               WEEK
     PATENT NO KIND DATE
                                           T.A
                                                PG
     ______
    WO 2001019862 A2 20010322 (200125) * EN 80
        RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC
            MW MZ NL OA PT SD SE SL SZ TZ UG ZW
         W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE
            DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG
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AU 2001013839 A 20010417 (200140) EP 1214339 A2 20020619 (200240)

YU ZA ZW

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

EN

KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

APPLICATION DETAILS:

PATENT NO K	IND	AP	PLICATION	DATE
WO 2001019862 AU 2001013839 EP 1214339		AU EP	2000-EP9034 2001-13839 2000-975853 2000-EP9034	20000914 20000914 20000914 20000914

FILING DETAILS:

-	TENT NO	KIND		PAT	ENT NO
	20010138			WO	200119862
•	1214339			WO	200119862

PRIORITY APPLN. INFO: GB 1999-22829 19990925; GB 1999-21693 19990914; GB 1999-21694 19990914

AN 2001-244783 [25] WPIDS

AB WO 200119862 A UPAB: 20010508

NOVELTY - Isolated Moraxella catarrhalis BASB129-BASB131 polypeptides (I) comprising a fully defined sequence of 344 (S2), 678 (S4), 469 (S6) amino acids, respectively as given in the specification, or an isolated polypeptide (Ia) which has 85% identity to (S2), (S4) or (S6), is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an \bar{i} mmunogenic fragment (II), of (I) which has the same \bar{i} mmunogenic activity as (I);
- (2) an isolated polynucleotide (III), or its complementary nucleotide sequence comprising a nucleotide sequence:
- (i) encoding a polypeptide that has 85% identity over the entire length of (S2), (S4), (S6);
- (ii) that has 85% identity over the entire length of the nucleotide sequence encoding region which encodes (S2), (S4), (S6);
- (iii) which has 85% identity over the entire length of a fully defined nucleotide sequence of 1035 (S1), 2037 (S3), 1410 (S5), base pairs as given in the specification;
- (iv) comprising a nucleotide sequence encoding (I) obtainable by screening an appropriate library under stringent hybridization conditions with a labeled probe having the sequence of (S1), (S3), (S5);
 - (v) encoding (S2), (S4) or (S6); or
 - (vi) an isolated polynucleotide comprising (S1), (S3) or (S5);
- (3) an expression vector (IV), or a recombinant live microorganism comprising (III);
- (4) a host cell (V) comprising (IV), or a subcellular fraction or membrane of the host cell expressing (I);
 - (5) preparation of (I) or (II);
- (6) expressing (III) involves transforming (V) with (IV) which contains any one of the polynucleotides (III) given above and culturing (V) under suitable conditions to express (III);
 - (7) a vaccine composition which comprises (I) or (II);
 - (8) a vaccine composition which comprises (III);
 - (9) an antibody (Ab) immunospecific for (I) or (II); and
- (10) a therapeutic composition comprising an antibody directed against (I) useful in treating humans with M.catarrhalis disease.

 ACTIVITY Antiinflammatory; auditory.

MECHANISM OF ACTION - Gene therapy; vaccine; initial physical

attraction between a pathogen and a mammalian extracellular matrix protein inhibitor.

The biological activity of (I) was tested in mice. Groups of mice were immunized with BASB129, BASB130 and BASB131 vaccine. After the booster, the mice were challenged by bacterial suspension into the nostril under anesthesia. Mice were killed between 30 minutes and 24 hours after challenge and the lungs were removed and homogenized. The log10 weighted mean number of colony forming unit (CFU)/lung was determined by counting the colonies grown on agar plates after plating of dilutions of the homogenate. The arithmetic mean of the log10 weighted mean number of CFU/lung and the standard deviations were calculated for each group. Results were analyzed statistically. Results showed that BASB129, BASB130 and BASB131 vaccine induced significant lung clearance as compared to the control group.

USE - The composition comprising (I), (II) or (III) is useful for preparation of a medicament used for generating an immune response in an animal. (I) is also useful as diagnostic reagent for M.catarrhalis which involves identifying (I), an antibody against (I) present within the biological sample from an animal suspected of having such an infection (claimed). Fragments of (I) are useful for producing corresponding full length polypeptides by peptide synthesis. The polynucleotides may be used as hybridization probes for RNA, cDNA and genomic DNA to isolate full-length cDNAs and genomic clones encoding BASB129-BASB131 and to isolate cDNA and genomic clones of other genes that have high sequence identity to BASB129-BASB131 gene. The polynucleotide sequences can also be used in the discovery and development of antibacterial compounds. The encoded protein can be used as target for the screening of antibacterial drugs. Additionally, the polynucleotide sequences encoding the amino terminal regions of the encoded protein or Shine-Dalgarno or other translation facilitating sequences of the respective mRNA can be used to construct antisense sequences to control the expression of the coding sequence of interest. The polynucleotides are also useful as diagnostic reagents in which the mutations in the polynucleotide sequence may be detected and used to diagnose and/or prognose the infection or its stage or course. The polynucleotides may be used as components of arrays which have diagnostic and prognostic uses. Antibodies against (I) are useful for treating bacterial infections and to isolate or identify clones expressing (I) or (II), to purify the polypeptides by affinity chromatography. The polynucleotides and polypeptides are used as research reagents and materials for discovery of treatments of and diagnostics for human diseases. The polynucleotides derived from (S1), (S3) or (S5) are used as PCR (polymerase chain reaction) primers. The polynucleotides are also useful in the diagnosis of the stage of infection and type of infection the pathogen has attained. The polypeptides and polynucleotides are used to block the initial physical interaction between a gram negative and/or gram positive bacteria to mammalian, host thus preventing the sequelae of infection. The polynucleotides encoding certain non-variable regions of bacterial cell surface protein are used in polynucleotide constructs which are useful for genetic immunization experiments in animal models of infection with M.catarrhalis to identify protein groups able to provoke a prophylactic or therapeutic immune response. The vaccine comprising (I), (II) or (III) is useful for treating Moraxella catarrhalis infections such as sinusitis, nosocomial infections, otitis media and pneumonia.

Dwg.0/0

L26 ANSWER 4 OF 20 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2003

2001-159876 [16] WPIDS

DOC. NO. NON-CPI: DOC. NO. CPI:

N2001-116486 C2001-047628

TITLE:

New BASB117 polypeptides from Moraxella catarrhalis strain MC2931 (ATCC 43617), useful as therapeutic agents or vaccines against bacterial (especially M.

catarrhalis) infections, e.g. otitis media or

pneumonia.

DERWENT CLASS:

B04 D16 S03 THONNARD, J

INVENTOR(S): PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

95

WO 2001009339 A2 20010208 (200116) * EN 79

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG

KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

AU 2000065688 A 20010219 (200129)

EP 1206547 A2 20020522 (200241) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	API	PLICATION	DATE
WO 2001009339 AU 2000065688 EP 1206547		AU EP	2000-EP7422 2000-65688 2000-953131 2000-EP7422	20000731 20000731 20000731 20000731

FILING DETAILS:

PATENT NO	KIND 		ENT NO
	B A Based on	WO	200109339 200109339

PRIORITY APPLN. INFO: GB 1999-18206 19990803

AN 2001-159876 [16] WPIDS

AB WO 200109339 A UPAB: 20010323

NOVELTY - Moraxella catarrhalis strain MC2931 (ATCC 43617) BASB117 polypeptides, both of 216 amino acids (I and II) as defined in the specification, are new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

(1) an isolated polypeptide (P1) comprising an amino acid sequence which has at least 85%, preferably 100%, identity to (I) or

(II) over their entire length;

- (2) an immunogenic fragment (P2) of the polypeptide, in which the immunogenic activity of the fragment is substantially the same as (I) or (II);
 - (3) an isolated polynucleotide (N1) selected from:
 - (a) a nucleotide sequence encoding (I), (II), P1 or P2;
- (b) an isolated polynucleotide comprising a nucleotide sequence encoding a polypeptide that has at least 85%, preferably 95%, identity to (I) or (II) over its entire length, or a nucleotide sequence complementary to the isolated polynucleotide;
- (c) an isolated polynucleotide comprising a nucleotide sequence that has at least 85%, preferably 95%, identity to a nucleotide sequence encoding (I) or (II) over the entire coding region, or a nucleotide sequence complementary to the isolated polynucleotide;

(d) an isolated polynucleotide comprising the 648 (III) or 651 basepair (bp) sequence (IV) fully defined in the specification;

- (e) an isolated polynucleotide comprising a nucleotide sequence which has at least 85%, preferably 95%, identity to (I) or (II) over its entire length, or a nucleotide sequence complementary to the isolated polynucleotide;
- (f) a nucleotide sequence encoding (I) or (II) obtainable by screening an appropriate library, under stringent conditions, with a labeled probe having the sequence of (III), (IV) or its fragments;
- (4) an expression vector or a recombinant live microorganism comprising N1;
- (5) a host cell comprising the expression vector of (4), or a subcellular fraction or membrane of the host cell expressing P1;
- (6) a process for producing (I), (II), P1 or P2 by culturing
 the host cell of (5);
- (7) a process for expressing N1 comprising transforming a host cell with the expression vector of (4) and culturing the host cell;
- (8) a vaccine compositions comprising (I), (II), P1 or P2 or N1;
 - (9) an antibody immunospecific for (I), (II), P1 or P2;
- (10) a method for diagnosing a Moraxella catarrhalis infection comprising identifying (I), (II), P1 or P2 or the antibody of (9) present within a biological sample from an animal suspected of having such an infection; and
- (11) a therapeutic composition for treating humans with Moraxella catarrhalis disease, comprising at least one antibody against (I), (II), P1 or P2.

ACTIVITY - Antibacterial; ophthalmological; antiinflammatory. MECHANISM OF ACTION - Vaccine; gene therapy.

Groups of mice were immunized with the polypeptide (BASB117) or with a killed whole cells (kwc) preparation of Moraxella catarrhalis or sham immunized.

After booster, the mice were challenged by instillation of bacterial suspension into the nostril under anaesthesia. Mice were killed between 30 minutes and 24 hours after challenge and the lungs were removed aseptically and homogenized individually. The log10 weighted mean number of colony forming units (CFU)/lung was determined by counting the colonies grown on agar plates after plating of dilutions of the homogenate. The arithmetic mean of the log10 weighted mean number of CFU/lung and the standard deviations were calculated for each group.

No results are given.

USE - The composition comprising an immunologic amount of the polypeptide or polynucleotide is useful for preparing a medicament

for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection (all claimed). The polypeptides may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptides or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the polypeptides or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging of the disease, determining the response of an infectious organism to drugs. Dwg.0/2

L26 ANSWER 5 OF 20 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2001-159875 [16] WPIDS

DOC. NO. NON-CPI: DOC. NO. CPI:

N2001-116485

TITLE:

C2001-047627 New BASB116 polypeptides from Moraxella catarrhalis

strain MC2931 (ATCC 43617), useful as therapeutic agents or vaccines against bacterial (especially M.

catarrhalis) infections, e.g. otitis media or

pneumonia.

DERWENT CLASS: INVENTOR(S):

B04 D16 S03 THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

PATENT INFORMATION:

PATENT NO	KIND	DATE	WEEK	LA	PG

95

WO 2001009338 A1 20010208 (200116) * EN 79

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW
W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE
DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG
KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ
PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

AU 2000062788 A 20010219 (200129)

EP 1206545 A1 20020522 (200241) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO KI	ND	APPLICATION	DATE
WO 2001009338 AU 2000062788 EP 1206545	A A1	WO 2000-EP7421 AU 2000-62788 EP 2000-949429 WO 2000-EP7421	20000731 20000731 20000731 20000731

FILING DETAILS:

PRIORITY APPLN. INFO: GB 1999-18279 19990803

AN 2001-159875 [16] WPIDS

AB WO 200109338 A UPAB: 20010323

NOVELTY - Two Moraxella catarrhalis strain MC2931 (ATCC 43617) BASB116 polypeptides, both of 98 amino acids (I and II) as defined in the specification, are new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated polypeptide (P1) comprising an amino acid sequence which has at least 85%, preferably 100%, identity to (I) or (II) over their entire length;
- (2) an immunogenic fragment (P2) of the polypeptide, in which the immunogenic activity of the fragment is substantially the same as (I) or (II);
 - (3) an isolated polynucleotide (N1) selected from:
 - (a) a nucleotide sequence encoding (I), (II), P1 or P2;
- (b) an isolated polynucleotide comprising a nucleotide sequence encoding a polypeptide that has at least 85% identity to (I) or (II) over its entire length, or a nucleotide sequence complementary to the isolated polynucleotide;
- (c) an isolated polynucleotide comprising a nucleotide sequence that has at least 85%, preferably 95%, identity to a nucleotide sequence encoding (I) or (II) over the entire coding region, or a nucleotide sequence complementary to the isolated polynucleotide;
- (d) an isolated polynucleotide comprising the 297 (III) or 294(IV) basepair (bp) sequence fully defined in the specification;
- (e) an isolated polynucleotide comprising a nucleotide sequence which has at least 85%, preferably 95%, identity to (I) or (II) over its entire length, or a nucleotide sequence complementary to the isolated polynucleotide;
- (f) a nucleotide sequence encoding (I) or (II) obtainable by screening an appropriate library, under stringent conditions, with a labeled probe having the sequence of (III), (IV) or its fragments;
- (4) an expression vector or a recombinant live microorganism comprising N1;
- (5) a host cell comprising the expression vector of (4), or a subcellular fraction or membrane of the host cell expressing Pl;
- (6) a process for producing (I), (II), P1 or P2 by culturing the host cell of (5);
- (7) a process for expressing N1 comprising transforming a host cell with the expression vector of (4) and culturing the host cell;
- (8) a vaccine compositions comprising (I), (II), P1 or P2 or N1;
 - (9) an antibody immunospecific for (I), (II), P1 or P2;
- (10) a method for diagnosing a Moraxella catarrhalis infection comprising identifying (I), (II), P1 or P2 or the antibody of (9) present within a biological sample from an animal suspected of having such an infection; and
- (11) a therapeutic composition for treating humans with Moraxella catarrhalis disease, comprising at least one antibody

against (I), (II), Pl or P2.

ACTIVITY - Antibacterial; ophthalmological; antiinflammatory.

MECHANISM OF ACTION - Vaccine; gene therapy.

Groups of mice were immunized with the polypeptide (BASB116) or with a killed whole cells (kwc) preparation of Moraxella catarrhalis or sham immunized.

After booster, the mice were challenged by instillation of bacterial suspension into the nostril under anaesthesia. Mice were killed between 30 minutes and 24 hours after challenge and the lungs were removed aseptically and homogenized individually. The log10 weighted mean number of colony forming units (CFU)/lung was determined by counting the colonies grown on agar plates after plating of dilutions of the homogenate. The arithmetic mean of the log10 weighted mean number of CFU/lung and the standard deviations were calculated for each group.

No results are given.

USE - The composition comprising an immunologic amount of the polypeptide or polynucleotide is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection (all claimed). The polypeptides may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptides or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the polypeptides or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging of the disease, determining the response of an infectious organism to drugs. Dwg.0/2

L26 ANSWER 6 OF 20 WPIDS. (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2001-159874 [16] WPIDS

DOC. NO. NON-CPI:

N2001-116484 C2001-047626

DOC. NO. CPI:

TITLE:

New BASB122 and BASB124 polypeptides and

polynucleotides from Moraxella catarrhalis strain

ATCC 43617, useful as therapeutic agents or

vaccines against bacterial infections, e.g. otitis

media or pneumonia.

DERWENT CLASS:

B04 D16 S03

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

95

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001009337 A2 20010208 (200116) * EN 75

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000065683 A 20010219 (200129) EP 1204749 A2 20020515 (200239) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO KI	ND	API	LICATION	DATE
WO 2001009337 AU 2000065683 EP 1204749		AU E.P.	2000-EP7365 2000-65683 2000-953120 2000-EP7365	20000731 20000731 20000731 20000731

FILING DETAILS:

PA'	rent no k	IND			PA'	PENT NO
AU	2000065683		Based	on	WO	200109337
EΡ	1204749	A2	Based	on	WO	200109337

PRIORITY APPLN. INFO: GB 1999-18036 19990730; GB 1999-18034 19990730

AN 2001-159874 [16] WPIDS

AB WO 200109337 A UPAB: 20010323

NOVELTY - New isolated polypeptides, comprising either of two 111 amino acid (I) or two 328 amino acid (II) sequences from Moraxella catarrhalis, all fully defined in the specification, or an at least 85 % identical sequence over their entire length, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated polynucleotide encoding the novel polypeptide, comprising:
 - (a) a sequence encoding the novel polypeptide;
- (b) a sequence having at least 85 % identity to (a) over its entire length;
- (c) a 336 (III) or 987 (IV) base pair sequence, both fully defined in the specification;
- (d) a sequence at least 85 % identical to (III) or (IV) over their entire length;
 - (e) the complements of (a)-(d); or
- (f) a sequence encoding (I) or (II) obtained by screening a library, under stringent conditions, with a labeled probe having (III), (IV), or fragments of them;
- (2) a statement vector or a recombinant live microorganism, comprising the polynucleotide of (1);
- (3) a host cell comprising the vector of (2), or a subcellular fraction or membrane of the host cell expressing the novel polypeptide;
- (4) a process for producing the novel polypeptide, comprising culturing the host cell of (3) under expression conditions, and recovering the polypeptide;
 - (5) a process for expressing the polynucleotide of (1),

comprising transforming a host cell with the vector of (2), and culturing the cell for expression of the polynucleotide;

- (6) a vaccine composition comprising the novel polypeptide or the polynucleotide of (1), and a carrier;
- (7) an antibody immunospecific for the novel polypeptide or its immunological fragment;
- (8) a method for diagnosing a M. catarrhalis infection, comprising identifying the novel polypeptide or the antibody of (7) present within a biological sample; and
- (9) a therapeutic composition comprising at least one antibody against the novel polypeptide.

ACTIVITY - Antibacterial; antiinflammatory; auditory. MECHANISM OF ACTION - Vaccine; gene therapy.

No biological data is given.

USE - The composition comprising an immunologic amount of the polypeptide or polynucleotide is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection. (All claimed). The polypeptides may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptides or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the polypeptides or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging of the disease, determining the response of an infectious organism to drugs. Dwg.0/0

L26 ANSWER 7 OF 20 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2001-159873 [16] WPIDS

DOC. NO. NON-CPI:

N2001-116483

DOC. NO. CPI:

C2001-047625

TITLE:

New BASB119 polypeptides and polynucleotides from Moraxella catarrhalis strain ATCC 43617, useful as therapeutic agents or vaccines against bacterial

infections, e.g. otitis media or pneumonia.

DERWENT CLASS:

B04 D16 S03

INVENTOR (S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

95

WO 2001009336 A1 20010208 (200116) * EN 82

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ T% UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ

PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000069887 A 20010219 (200129)

EP 1206549 A1 20020522 (200241) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	API	PLICATION	DATE
WO 2001009336 AU 2000069887 EP 1206549		AU EP	2000-EP7363 2000-69887 2000-958324 2000-EP7363	20000731 20000731 20000731 20000731

FILING DETAILS:

PAT	rent no 'K				PAT	ENT NO
AU	2000069887				WO	200109336
EΡ	1206549	A1	Based	on	WO	200109336

PRIORITY APPLN. INFO: GB 1999-18302 19990803

AN 2001-159873 [16] WPIDS

AB WO 200109336 A UPAB: 20010323

NOVELTY - New isolated polypeptides, comprising either of two 171 residue amino acid sequences (I and II) from Moraxella catarrhalis, both fully defined in the specification, or a sequence at least 85 % identical to (I) or (II), over their entire length, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated polynucleotide encoding the novel polypeptide, comprising:
 - (a) a sequence encoding (I) or (II);
- (b) a sequence having at least 85 % identity to the sequence encoding (I) or (II) over the entire coding region;
- (c) a 516 (III) or 513 (IV) base pair sequence, fully defined in the specification;
- (d) a sequence having at least 85 % identity to (III) or (IV) over their entire length;
 - (e) the complements of (a)-(d); or
- (f) a sequence encoding (I) or (II) obtained by screening a library, under stringent conditions, with a labeled probe having (III), (IV), or fragments of (III) or (IV);
- (2) an statement vector or a recombinant live microorganism comprising the polynucleotide of (1);
- (3) a host cell comprising the vector of (2), or a subcellular fraction or membrane of the host cell expressing the novel polypeptide;
- (4) a process for producing the novel polypeptide, comprising culturing the cell of (3) under expression conditions, and recovering the polypeptide;
- (5) a process for expressing the polynucleotide of (1), comprising transforming a host cell with the vector of (2), and culturing the host cell for expression of the polynucleotide;
- (6) vaccine compositions comprising the novel polypeptide or the polynucleotide of (1), and a carrier;

- (7) an antibody immunospecific for the novel polypeptide or its immunological fragment;
- (8) a method for diagnosing a M. catarrhalis infection, comprising identifying the novel polypeptide or the antibody present within a biological sample; and
- (9) a therapeutic composition comprising at least one antibody against the novel polypeptide.

ACTIVITY - Antibacterial; antiinflammatory; auditory.

MECHANISM OF ACTION - Vaccine; gene therapy.

Groups of mice were immunized either with the polypeptide (BASB119) adsorbed onto AlPO4 (10 micro g BASB119 onto 100 micro g of AlPO4), with a killed whole cell (kwc) preparation of M. catarrhalis strain ATCC 43617 adsorbed onto AlPO4, or with 100 micro g AlPO4 without antigen. The mice were challenged with 5 multiply 105 colony forming units (CFU) of live M. catarrhalis strain ATCC 43617 bacteria. The log10 weighted mean number of CFU/lung and the standard deviation 4 hours after challenge was calculated for each group. Sham immunized mice had 5.41 (+/-0.2) log10 CFU/lungs 4 hours after challenge. The kwc preparation induced significant lung clearance as compared to the control group (1.58 log difference). BASB119 vaccine induced a 1.34 log difference in lung clearance, which was significantly different from the control.

USE - The composition comprising the novel polypeptide or polynucleotide is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection. (All claimed). The polypeptides may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptides or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the polypeptides or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging of the disease, determining the response of an infectious organism to drugs. Dwg.0/3

L26 ANSWER 8 OF 20 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2001-159872 [16] WPIDS

DOC. NO. NON-CPI: DOC. NO. CPI:

N2001-116482

DOC. NO

C2001-047624

TITLE:

New BASB120 polypeptides and polynucleotides from Moraxella catarrhalis strain American Type Culture Collection 43617, for use as therapeutic agents or vaccines against bacterial infections, e.g. otitis media or pneumonia.

DERWENT CLASS:

B04 D16 S03

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

. . .

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA WO 2001009335 A2 20010208 (200116) * EN 75 RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW AU 2000064397 A 20010219 (200129) A2 20020522 (200241) EN EP 1206546 R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI APPLICATION DETAILS: KIND APPLICATION PATENT NO WO 2001009335 A2 WO 2000-EP7361 20000731 AU 2000-64397 AU 2000064397 A 20000731 EP 1206546 A2 EP 2000-951472 20000731 WO 2000-EP7361 20000731 FILING DETAILS: PATENT NO KIND PATENT NO AU 2000064397 A Based on WO 200109335 EP 1206546 A2 Based on WO 200109335 PRIORITY APPLN. INFO: GB 1999-18281 19990803 2001-159872 [16] WPIDS WO 200109335 A UPAB: 20010323 NOVELTY - An isolated polypeptide (PP) comprising: (a) a sequence of 250 amino acids (I) from Moraxella catarrhalis, given in the specification; or (b) an amino acid sequence, which has at least 85% identity to (I), over the entire length of (I), is new. DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following: (1) an immunogenic fragment of the polypeptide, in which the

(i) a nucleotide sequence encoding (PP);

immunogenic activity of the fragment is the same as (I);

AN

comprising:

(ii) a nucleotide sequence having 85% identity to the nucleotide sequence encoding (I) over the entire coding region;

(iii) a 753 base pair (bp) DNA sequence (II), given in the specification;

(iv) a nucleotide sequence having 85% identity to (II) over the entire length of (II);

(2) isolated polynucleotides, which encode the polypeptides,

(v) the complements of (i)-(iv); or

(vi) a nucleotide sequence encoding (I) obtainable by screening an appropriate library, under stringent conditions, with a labeled probe having (II) or its fragments;

(3) an expression vector or a recombinant live microorganism

comprising (2);

(4) a host cell comprising the expression vector, or a subcellular fraction or membrane of the host cell expressing (PP); (5) producing (PP) comprising culturing (4) to produce (PP) and

recovering (PP) from the culture medium;

(6) expressing (2) comprising transforming a host cell with the expression vector and culturing the host cell for expression of any of the polynucleotides;

(7) vaccine compositions comprising (PP) or (2), and a pharmaceutical carrier;

(8) an antibody immunospecific for (PP) or immunological fragment of (1);

(9) diagnosing a M. catarrhalis infection comprising identifying (PP) or the antibody of (8) present within a biological sample from an animal suspected of having such an infection;

(10) using the compositions of (7) for preparing a medicament for use in generating an immune response in an animal; and

(11) a therapeutic composition comprising the antibody of (8). ACTIVITY - Antibacterial; antiinflammatory; pulmonary. MECHANISM OF ACTION - Vaccine; gene therapy. Clinical test

details are described but no results are given.

USE - A composition comprising an immunologic amount of (PP) or a polynucleotide encoding it, is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection (all claimed). The polypeptides may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptides or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the polypeptides or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging diseases, and determining the response of an infectious organism to drugs. Dwg.0/2

L26 ANSWER 9 OF 20 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2001-159871 [16] WPIDS

DOC. NO. NON-CPI:

N2001-116481

DOC. NO. CPI:

C2001-047623

TITLE:

New BASB118 polypeptides and polynucleotides from Moraxella catarrhalis strain American Type Culture Collection 43617, for use as therapeutic agents or vaccines against bacterial infections, e.g. otitis

media or pneumonia.

DERWENT CLASS:

B04 D16 S03

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

95

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001009334 A1 20010208 (200116) * EN 77

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE

DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ

PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000068330 A 20010219 (200129)

EP 1206548 A1 20020522 (200241) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	AP	PLICATION	DATE
WO 2001009334 AU 2000068330 EP 1206548	•	AU EP	2000-EP7360 2000-68330 2000-956353 2000-EP7360	20000731 20000731 20000731 20000731

FILING DETAILS:

PA	TENT	NO	KIND		•	PA	TENT	NO
AU	2000	006833	0 A	Based	on	WO	2001	.09334
ΕP	1206	6548	A1	Based	on	WO	2001	09334

PRIORITY APPLN. INFO: GB 1999-18208 19990803

AN 2001-159871 [16] WPIDS

AB WO 200109334 A UPAB: 20010323

NOVELTY - An isolated polypeptide comprising:

- (a) a sequence of 386 amino acids (I) from Moraxella catarrhalis, given in the specification; or
- (b) an amino acid sequence, which has 85% identity to (I), over the entire length of (I), is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an immunogenic fragment of the new polypeptide, in which the immunogenic activity of the fragment is the same as (I);
- (2) isolated polynucleotides, which encode the new polypeptide, comprising:
 - (i) a nucleotide sequence encoding (a) or (b);
- (ii) a nucleotide sequence that has 85% identity to the nucleotide sequence encoding (I) over the entire coding region;
- (iii) a 1161 base pair (bp) DNA sequence (II), given in the specification;
- (iv) a nucleotide sequence that has 85% identity to (II) over the entire length of (II);
 - (v) the complements of (i)-(iv); or
- (vi) a nucleotide sequence encoding (I) obtainable by screening an appropriate library, under stringent conditions, with a labeled probe having (II) or its fragments;
- (3) an expression vector or a recombinant live microorganism comprising an isolated polynucleotide of (2);

- (4) a host cell comprising the expression vector of (3), or a subcellular fraction or membrane of the host cell expressing the new polypeptide;
- (5) producing the new polypeptide comprising culturing (4) to produce the new polypeptide and recovering it from the culture medium;
- (6) expressing a polynucleotide of (2) comprising transforming a host cell with the expression vector of (3) and culturing the host cell for expression of any of the polynucleotides;
- (7) vaccine compositions comprising the new polypeptide or polynucleotide of (2), and a pharmaceutical carrier;
- (8) an antibody immunospecific for the new polypeptide or immunological fragment;
- (9) diagnosing a M. catarrhalis infection comprising identifying the new polypeptide or the antibody of (8) present within a biological sample from an animal suspected of having such an infection; and
 - (10) a therapeutic composition comprising an antibody of (8). ACTIVITY Antibacterial; antiinflammatory; pulmonary.

MECHANISM OF ACTION - Vaccine; gene therapy. Groups of mice were immunized either with the polypeptide (BASB118) adsorbed onto AlPO4 (10 micro g BASB118 onto 100 micro g of AlPO4), with a killed whole cell (kwc) preparation of M. catarrhalis strain American type Culture Collection (ATCC) 43617 adsorbed onto AlPO4, or with 100 micro g AlPO4 without antigen. The mice were challenged with 5 multiply 105 colony forming units (CFU) of live M. catarrhalis strain ATCC 43617 bacteria. The log10 weighted mean number of CFU/lung and the standard deviation 4 hours after challenge was calculated for each group. Sham immunized mice had 5.66 (+/-0.18) log10 CFU/lungs 4 hours after challenge. The kwc preparation induced significant lung clearance as compared to the control group (1.3 log difference). BASB118 vaccine induced a 0.43 log difference in lung clearance, which was significantly different from the control.

USE - A composition comprising an immunologic amount of the new polypeptide or polynucleotide encoding it, is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection (all claimed). The polypeptide may also be used as a prophylactic agent of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptides or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the new polypeptide or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging diseases, and determining the response of an infectious organism to drugs. Dwg.0/1

L26 ANSWER 10 OF 20 WPIDS (C) 2002 THOMSON DERWENT ACCESSION NUMBER: 2001-159870 [16] WPIDS

DOC. NO. NON-CPI:

N2001-116480

DOC. NO. CPI:

C2001-047622

TITLE:

New BASB123 polypeptides and polynucleotides from Moraxella catarrhalis strain American type Culture Collection 43617, for use as therapeutic agents or vaccines against bacterial infections, e.g. otitis

media or pneumonia.

DERWENT CLASS:

B04 D16 S03 THONNARD, J

INVENTOR(S):
PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

94

WO 2001009333 A2 20010208 (200116) * EN 79

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE

DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG

KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ

PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

AU 2000069880 A 20010219 (200129)

APPLICATION DETAILS:

PATENT NO KIND	APPLICATION	DATE
WO 2001009333 A2	WO 2000-EP7296	20000727
AU 2000069880 A	AU 2000-69880	20000727

FILING DETAILS:

PATENT NO	KIND			PAT	ENT	NO
AU 2000069	880 A	Based	on	WO	2001	109333

PRIORITY APPLN. INFO: GB 1999-17975 19990730

AN 2001-159870 [16] WPIDS

AB WO 200109333 A UPAB: 20010323

NOVELTY - An isolated polypeptide comprising:

(a) a sequence comprising one of two 167 amino acid sequences (designated I and II) from Moraxella catarrhalis, given in the specification; or

(b) an amino acid sequence, which has 85% identity to (I) or (II), over the entire length of (I) or (II), respectively, is new. DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

(1) an immunogenic fragment of the new polypeptide, in which the immunogenic activity of the fragment is the same as (I) or (II);

(2) isolated polynucleotides, which encode the new polypeptide, comprising:

(i) a nucleotide sequence encoding (a) or (b);

(ii) a nucleotide sequence that has 85% identity to the nucleotide sequence encoding (I) or (II) over the entire coding region;

(iii) a 504 base pair (bp) (III) or 501 bp (IV) DNA sequence,

given in the specification;

(iv) a nucleotide sequence that has 85% identity to (III) or (IV) over the entire length of (III) or (IV), respectively;

(v) the complements of (i)-(iv); or

(vi) a nucleotide sequence encoding (I) or (II) obtainable by screening an appropriate library, under stringent conditions, with a labeled probe having (III), (IV), or fragments of (III) or (IV);

(3) an expression vector or a recombinant live microorganism comprising a polynucleotide of (2);

(4) a host cell comprising the expression vector of (3), or a subcellular fraction or membrane of the host cell expressing the new polypeptide;

(5) producing the new polypeptide comprising culturing (4) t produce the polypeptide and recovering it from the culture medium;

- (6) expressing a polynucleotide of (2) comprising transforming a host cell with the expression vector of (3) and culturing the host cell for expression of any of the polynucleotides;
- (7) vaccine compositions comprising the new polypeptide or polynucleotide of (2), and a pharmaceutical carrier;
- (8) an antibody immunospecific for the new polypeptide or an immunological fragment;
- (9) diagnosing a M. catarrhalis infection comprising identifying the new polypeptide or the antibody of (8) present within a biological sample from an animal suspected of having such an infection; and
 - (10) a therapeutic composition comprising an antibody of (8). ACTIVITY Antibacterial.

MECHANISM OF ACTION - Vaccine; gene therapy. Clinical details are described but no results are given.

USE - A composition comprising an immunologic amount of the new polypeptide or polynucleotide encoding it, is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection (all claimed). The polypeptides may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptide or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the polypeptide or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging of diseases, and determining the response of an infectious organism to drugs. Dwq.0/2

L26 ANSWER 11 OF 20 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2001-159869 [16] WPIDS

DOC. NO. NON-CPI: N2001-116479 DOC. NO. CPI: C2001-047621

TITLE:

New BASB115 polypeptide from Moraxella catarrhalis strain MC2931 (ATCC 43617), useful as a therapeutic agent or vaccine against bacterial

(especially M. catarrhalis) infections, e.g. otitis media or pneumonia.

DERWENT CLASS:

B04 D16 S03

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

95

PATENT INFORMATION:

PATENT 1	NO KIND	DATE	WEEK	LA	PG

WO 2001009332 A2 20010208 (200116)* EN 72

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

AU 2000068323 A 20010219 (200129)

EP 1204752 A2 20020515 (200239) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	AP	PLICATION	DATE
WO 2001009332 AU 2000068323 EP 1204752		AU EP	2000-EP7294 2000-68323 2000-956339 2000-EP7294	20000727 20000727 20000727 20000727

FILING DETAILS:

PATENT NO	KIND	PATENT NO
AU 200006832	3 A Based on	WO 200109332
EP 1204752	A2 Based on	WO 200109332

PRIORITY APPLN. INFO: GB 1999-18003

19990730

AN 2001-159869 [16] WPIDS

AB WO 200109332 A UPAB: 20010323

NOVELTY - A Moraxella catarrhalis strain MC2931 (ATCC 43617) BASB115 polypeptide of 199 amino acids (I) as defined in the specification, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated polypeptide (P1) comprising an amino acid sequence which has at least 85%, preferably 100%, identity to (I) over its entire length;
- (2) an immunogenic fragment (P2) of the polypeptide, in which the immunogenic activity of the fragment is substantially the same as (I);
 - (3) an isolated polynucleotide (N1) selected from:
 - (a) a nucleotide sequence encoding (I), P1 or P2;
- (b) an isolated polynucleotide comprising a nucleotide sequence encoding a polypeptide that has at least 85%, preferably 95%, identity to (I) over its entire length, or a nucleotide sequence

complementary to the isolated polynucleotide;

- (c) an isolated polynucleotide comprising a nucleotide sequence that has at least 85%, preferably 95%, identity to a nucleotide sequence encoding (I) over the entire coding region, or a nucleotide sequence complementary to the isolated polynucleotide;
- (d) an isolated polynucleotide comprising the 600 basepair (bp) sequence (II) fully defined in the specification;
- (e) an isolated polynucleotide comprising a nucleotide sequence which has at least 85%, preferably 95%, identity to (I) over its entire length, or a nucleotide sequence complementary to the isolated polynucleotide;
- (f) a nucleotide sequence encoding (I) obtainable by screening an appropriate library, under stringent conditions, with a labeled probe having the sequence of (II) or its fragments;
- (4) an expression vector or a recombinant live microorganism comprising N1;
- (5) a host cell comprising the expression vector of (4), or a subcellular fraction or membrane of the host cell expressing P1;
- (6) a process for producing (I), P1 or P2 by culturing the host cell of (5);
- (7) a process for expressing N1 comprising transforming a host cell with the expression vector of (4) and culturing the host cell;
 - (8) a vaccine compositions comprising (I), P1 or P2 or N1;
 - (9) an antibody immunospecific for (I), P1 or P2;
- (10) a method for diagnosing a M. catarrhalis infection comprising identifying (I), P1 or P2 or the antibody of (9) present within a biological sample from an animal suspected of having such an infection; and
- (11) a therapeutic composition for treating humans with M. catarrhalis disease, comprising at least one antibody against (I), P_1 or P_2

ACTIVITY - Antibacterial; ophthalmological; antiinflammatory. MECHANISM OF ACTION - Vaccine; gene therapy.

Groups of mice were immunized either with the polypeptide (BASB115) adsorbed onto AlPO4 (10 mu g BASB115 onto 100 mu g of AlPO4), with a killed whole cells (kwc) preparation of M. catarrhalis strain ATCC 43617 adsorbed onto AlPO4, or with 100 mu g AlPO4 without antigen. The mice were challenged with 5 x 105 colony forming units (CFU) of live M. catarrhalis strain ATCC 43617 bacteria. The log10 weighted mean number of CFU/lung and the standard deviation 4 hours after challenge was calculated for each group. Sham immunized mice had 5.66 (+/-0.18) log10 CFU/lungs 4 hours after challenge. The kwc preparation induced significant lung clearance as compared to the control group (1.76 log difference). BASB115 vaccine induced a 0.46 log difference in lung clearance, which was significantly different from the control.

USE - The composition comprising an immunologic amount of the polypeptide or polynucleotide is useful for preparing a medicament for generating an immune response in an animal. The therapeutic composition is useful in treating humans with M. catarrhalis infection (all claimed). The polypeptides may also be used as prophylactic agents of bacterial infections, particularly M. catarrhalis infections in mammals, especially humans. The polynucleotides are useful in therapy or prophylaxis, particularly genetic immunization against these infections or diseases. These diseases include otitis media in infants or children, pneumonia in elderlies, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in the

middle ear, infection of the upper respiratory tract, or inflammation of the middle ear. The polypeptides or polynucleotides may also be employed as research reagents and materials for discovering treatments of and diagnostics for diseases, particularly human diseases. In particular, the polypeptides or polynucleotides are useful in the discovery and development of antibacterial compounds, or for diagnosing diseases, staging of the disease, determining the response of an infectious organism to drugs. Dwg.0/1

L26 ANSWER 12 OF 20 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2001-168707 [17] WPIDS

DOC. NO. NON-CPI:

N2001-121639

DOC. NO. CPI:

C2001-050432

TITLE:

New BASB125 polypeptide isolated from Moraxella catarrhalis for treating, preventing and diagnosing diseases associated with M. catarrhalis infection

in mammals, e.g. otitis media in humans.

DERWENT CLASS:

B04 D16 S03

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

95

PATENT INFORMATION:

PATENT NO	KIND DATE	WEEK	LA PG	į
				_

WO 2001009331 A2 20010208 (200117) * EN 73

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

AU 2000064393 A 20010219 (200129)

A2 20020612 (200239) EP 1212424 EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	API	PLICATION	DATE
WO 2001009331 AU 2000064393 EP 1212424	*	AU EP	2000-EP7291 2000-64393 2000-951466 2000-EP7291	20000727 20000727 20000727 20000727

FILING DETAILS:

	IND	PATENT NO
AU 2000064393 EP 1212424	A Based on	WO 200109331 WO 200109331

PRIORITY APPLN. INFO: GB 1999-18041

19990730

2001-168707 [17] WPIDS AN

WO 200109331 A UPAB: 20010328 AB

> 308-4994 Searcher : Shears

NOVELTY - An isolated polypeptide having at least 85 % identity to a sequence (I) of 134 amino acids for a Moraxella catarrhalis BASB125 polypeptide, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for

the following:

(1) an isolated polypeptide of sequence (I);

(2) immunogenic fragments of the polypeptide having the same immunogenic activity as sequence (I);

(3) an isolated polynucleotide:

(i) having 85 % identity to a polynucleotide encoding the polypeptide, especially 85 % identity to sequence (II) of 405 base pairs (bp) encoding sequence (I);

(ii) complementary to a polynucleotide of (i);

(iii) encoding the new polypeptide; and

(iv) encoding sequence (I) and obtained by screening a library under stringent conditions using sequence (II) or a fragment as a probe;

(4) vectors or recombinant live microorganisms comprising the polynucleotide;

(5) host cells comprising the vector and subcellular fragments/membranes of the host cells expressing the polypeptide;

- (6) producing the new polypeptide comprising culturing the host cell of (5) to produce the polypeptide and recovering the polypeptide from the culture medium;
- (7) expressing (3) comprising transforming a host cell with an expression vector of (4) and culturing the host cell to express the polynucleotide;
 - (8) vaccine compositions comprising the new polypeptide or (3);

(9) antibodies specific for the new polypeptide, or immunological fragments of (2);

(10) diagnosing a M. catarrhalis infection comprising identifying the new polypeptide or an antibody immunospecific for the polypeptide, present within a biological sample from an animal suspected of having the infection;

(11) preparing a medicament for generating an immune response in an animal using a composition comprising the new polypeptide or (3); and

(12) a therapeutic composition for treating humans with M.catarrhalis disease comprising an antibody against the new polypeptide.

ACTIVITY - Antibacterial. A sequence (II) of 405 base pairs (bp) was isolated from M. catarrhalis strain American Type Culture Collection (ATCC) 43617 by standard molecular biological techniques a sequence (I) of 134 amino acids deduced. Mice were immunized with a BASB125 vaccine or a killed whole cell (kwc) M. catarrhalis preparation, or were sham immunized. After a booster, mice were challenged by instillation of bacterial suspension into the nostril under anaesthesia. Mice were killed 30 minutes-24 hours after challenge and lungs removed aseptically and homogenized. Homogenates were diluted and plated onto agar plates, and log10 weighted mean number of colony forming units/lung determined by counting. BASB125 vaccine and kwc preparations induced significant lung clearance of M. catarrhalis versus controls. No experimental data is given.

MECHANISM OF ACTION - Vaccine; gene therapy.

USE - The polypeptide, immunogenic fragments of the polypeptide, fusion proteins of the polypeptide, or polynucleotides encoding the polypeptide are used in vaccine compositions (claimed), optionally with another M. catarrhalis

antigen (claimed). They can also be included in medicaments for use in generating an immune response in an animal (claimed). The vaccines and medicaments are useful in preventing and/or treating microbial diseases, especially diseases associated with M. catarrhalis infection in mammals (especially humans). The polypeptides/polynucleotides may be used to produce antibodies, which can be used in compositions useful therapeutically to treat humans with M. catarrhalis diseases (claimed). M. catarrhalis is a Gram-negative bacteria frequently isolated from the human upper respiratory tract and responsible for several pathologies in humans e.g. otitis media in children, pneumonia, sinusitis etc. The polypeptides, polynucleotides and antibodies are also useful diagnostically e.g. in the detection of the polypeptides/antibodies in a biological sample from an animal to diagnose M. catarrhalis infection (claimed). The diagnostic assays are useful e.g. to detect diseases, determine the stage and type of infection, determine the effect of drugs etc. The polypeptides and polynucleotides can also be used to detect antagonists and agonists useful e.g. in preventing, inhibiting and/or treating disease. The polynucleotides are also useful in producing hybridization probes to isolate sequences encoding BASB125 and similar sequences. Dwg.0/0

L26 ANSWER 13 OF 20 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2001-182955 [18] WPIDS

DOC. NO. NON-CPI:

N2001-130566

DOC. NO. CPI:

C2001-054636

TITLE:

New BASB126 polypeptides of Moraxella catarrhalis useful for diagnostic, prophylactic and therapeutic purposes against microbial diseases, preferably

bacterial infections.

DERWENT CLASS:

B04 D16 S03

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

PATENT INFORMATION:

PATENT NO	KIND DATE	WEEK	LA	PG

95

WO 2001009329 A1 20010208 (200118) * EN 86

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ T7 UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

20010219 (200129) AU 2000068316 A

A1 20020515 (200239) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL RO SI

APPLICATION DETAILS:

PATENT NO KIND	APPLICATION.	DATE
WO 2001009329 A1	WO 2000-EP7280	20000727
AU 2000068316 A	AU 2000-68316	20000727

308-4994 Searcher : Shears

EP 1204750 A1

EP 2000-956332 20000727 WO 2000-EP7280 20000727

FILING DETAILS:

PATENT NO KI	IND	PATENT NO
AU 2000068316	A Based on	WO 200109329 WO 200109329

PRIORITY APPLN. INFO: GB 1999-18038 19990730

AN 2001-182955 [18] WPIDS

AB WO 200109329 A UPAB: 20010402

NOVELTY - An isolated BASB126 polypeptide (I) of Moraxella catarrhalis, comprises a sequence having at least 85% identity (over the entire length) to one of the two 192 amino acids sequences given in the specification.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an immunogenic fragment (II) of (I), where (II) has the same immunogenicity of (I);
 - (2) an isolated polynucleotide (III) encoding (I) (II);
- (3) an expression vector (IV) or a recombinant live microorganism, comprising (III);
- (4) a host cell (V) comprising (IV), or a subcellular fraction or membrane of (V) expressing (I);
- (5) producing (I) comprising culturing (V) and recovering the polypeptide from the culture medium;
- (6) expressing (III) comprising transforming (V) with (IV) and culturing under conditions sufficient for its expression;
 - (7) a vaccine (VI) comprising (I), (II) or (III);
 - (8) an antibody (VII) immunospecific for (I) or (II);
- (9) diagnosing Moraxella catarrhalis infection comprising identifying (I) or (VII) in a biological sample from an animal suspected of having such an infection; and
- (10) a therapeutic composition (VIII) for treating Moraxella catarrhalis infection comprising at least one (VII).

ACTIVITY - Antibacterial; antimicrobial; auditory; antiinflammatory.

MECHANISM OF ACTION - Vaccine.

Experimental protocols are described but no results are given. USE - (VI) is useful for preparing a medicament for use in generating immune response in an animal (claimed). (VIII) is useful for treating humans with Moraxella catarrhalis disease (claimed).

(I) and (III) are useful in the prevention, treatment and diagnosis of microbial diseases, preferably bacterial infections such as otitis media, pneumonia, sinusitis, nosocomial infections, and invasive diseases. (I) and (III) are useful as immunogens to produce antibodies, and to asses the binding of small molecule substrate and ligands in, for e.g., cells, cell-free preparations, chemical libraries and natural product mixtures. (I), (III) and (VII) are useful to configured screening methods for detecting the effect of added compounds and production of mRNA and/or polypeptides in the cells.

(III) is useful as a hybridization probe for RNA, cDNA and genomic DNA to isolate full-length cDNAs and genomic clones encoding BASB126 and to isolate cDNA and genomic clones of other genes that have a high identity particularly high sequence identity, to the

BASB126 gene. (II) has utility in diagnosis of the stage and type of infection, and also for therapeutic or prophylactic purposes, in particular genetic immunization. (II) is useful as a component of polynucleotide arrays, preferably high density arrays or grid. Dwg.0/4

L26 ANSWER 14 OF 20 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER:

2001-112459 [12] WPIDS

DOC. NO. NON-CPI:

N2001-082527

DOC. NO. CPI:

C2001-033488

TITLE:

Novel BASB110 polypeptides of Moraxella

catarrhalis, useful as a vaccine for treating

Moraxella catarrhalis infections.

DERWENT CLASS:

B04 D16 S03

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT: 95

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001000838 A1 20010104 (200112) * EN 88

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG

YU ZA ZW

AU 2000059779 A 20010131 (200124)

EP 1196589 A1 20020417 (200233) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO KIND	APPLICATION	DATE
WO 2001000838 A1 AU 2000059779 A EP 1196589 A1	WO 2000-EP5854 AU 2000-59779 EP 2000-945812 WO 2000-EP5854	20000623 20000623 20000623 20000623

FILING DETAILS:

PATENT NO	KIND	PAT	TENT NO
AU 200005977	9 A Based o	on WO	200100838
EP 1196589	Al Based o	on WO	200100838

PRIORITY APPLN. INFO: GB 1999-15031 19990625

AN 2001-112459 [12] WPIDS

AB WO 200100838 A UPAB: 20010302

NOVELTY - Isolated BASB110 polypeptides (I) of Moraxella catarrhalis, are new. The BASB110 polypeptide has the 322 (P1) or another 322 (P2) amino acid sequence defined in the specification.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated polypeptide (Ia) comprising an amino acid sequence which is at least 85%, preferably 95%, most preferably 100%, identical to the sequence, over its entire length, of P1 or P2;
- (2) an immunogenic fragment (Ib) of (I) or (Ia), where the activity of the fragment is substantially the same as P1 or P2;
 - (3) an isolated polynucleotide (II) encoding (I), (Ia) or (Ib);
- (4) an isolated polynucleotide (IIa) comprising a sequence encoding (Ia) or its complementary sequence;
- (5) an isolated polynucleotide (IIb) comprising a sequence having at least 85%, preferably 95%, most preferably 100% identity to a sequence encoding Pl or P2 over the entire coding region, or a nucleotide sequence complementary to the isolated polynucleotide;
- (6) an isolated polynucleotide (IIc) comprising a sequence having at least 85%, preferably 95%, most preferably 100% identical to the 969 (N1) or 966 (N2) nucleotides fully defined in the specification, or its complement;
- (7) an isolated polynucleotide (IId) comprising a sequence encoding P1 or P2, obtainable by screening an appropriate library under stringent hybridization conditions with labeled probe having the sequence of N1 or N2;
- (8) an expression vector (III) of a recombinant live microorganism, comprising (II), (IIa), (IIb), (IIc) or (IId);
- (9) a host cell (IV) comprising (III), or a subcellular fraction or membrane of (IV) expressing (Ia);
- (10) a process for producing (I), (Ia) or (Ib) comprising culturing (IV);
- (11) a process for expressing (II), (IIa), (IIb), (IIc) or (IId), comprising transforming (IV) with (III) and culturing transformed (IV) under conditions sufficient for its expression;
- (12) a vaccine composition (V) comprising (I), (Ia) or (Ib), or
 (II), (IIa), (IIb), (IIc) or (IId);
- (13) an antibody (Ab1) immunospecific for (I), (Ia) or (Ib); and
- (14) a method for diagnosing Moraxella catarrhalis infection, by identifying (I)-(Ib) or Abl present within a biological sample from an animal suspected of having such an infection.

ACTIVITY - Antibacterial.

MECHANISM OF ACTION - Vaccine.

Groups of mice are immunized with BASB110 vaccine. After the booster, the mice were challenged by instillation of bacterial suspension into the nostril under anaesthesia. Mice were killed between 30 minutes and 24 hours after challenge and the lungs were removed aseptically and homogenized individually. The log 10 weighted mean number of colony forming units (CFU)/lung was determined by counting the colonies grown on agar plates after plating of dilutions of the homogenate. The arithmetic mean of the log 10 weighted mean number of CFU/lung and the standard deviations were calculated for each group. Results were not given in the specification.

USE - The vaccine is useful for preparing a medicament for use in generating immune response in an animal (claimed). Abl is useful for treating humans suffering from Moraxella catarrhalis disease (claimed).

Polynucleotides encoding the BASB110 polypeptides have utility in diagnosis of the stage and type of infection, and also for therapeutic or prophylactic purposes, in particular genetic immunization.

WPIDS

Dwg.0/3

L26 ANSWER 15 OF 20 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2001-112458 [12]

DOC. NO. NON-CPI: N2001-082526 DOC. NO. CPI: C2001-033487

TITLE: New BASB113 polypeptide isolated from Moraxella catarrhalis bacterium, useful for diagnosing and

producing vaccines against bacterial infections

such as otitis media and pneumonia.

DERWENT CLASS: B04 D16 S03 INVENTOR(S): THONNARD, J

PATENT ASSIGNEE(S): (SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT: 95

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001000836 A1 20010104 (200112)* EN 86

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE

DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ

PL PT RO RU SD SE SG SI SK SI TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000059778 A 20010131 (200124)

EP 1196588 A1 20020417 (200233) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	API	PLICATION	DATE
WO 2001000836 AU 2000059778 EP 1196588		AU EP	2000-EP5851 2000-59778 2000-945811 2000-EP5851	20000623 20000623 20000623 20000623

FILING DETAILS:

PATENT NO K	IND	PATENT NO
AU 2000059778 EP 1196588		WO 200100836 WO 200100836

PRIORITY APPLN. INFO: GB 1999-15044 19990625

AN 2001-112458 [12] WPIDS

AB WO 200100836 A UPAB: 20010302

NOVELTY - An isolated polypeptide (I) comprising an amino acid sequence which has 85% identity to the Moraxella catarrhalis BASB113 polypeptide sequence of 224 (S2) or 224 (S4) amino acids respectively as given in the specification, or has a sequence of (S2) or (S4), is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

(1) an immunogenic fragment (II) of (I) which has the same

immunogenic activity as (I);

- (2) an isolated polynucleotide (III), or its complementary nucleotide sequence comprising a nucleotide sequence:
- (i) encoding a polypeptide that has 85% identity over the entire length of (S2) or (S4);
- (ii) that has 85% identity over the entire length of the nucleotide sequence encoding region which encodes (S2) or (S4);
- (iii) which has 85% identity over the entire length of a fully defined nucleotide sequence of 675 (S1) or 672 (S3) base pairs as given in the specification; and
- (iv) comprising a nucleotide sequencing encoding (I) obtainable by screening an appropriate library under stringent hybridization conditions with a labeled probe with the sequence of (S1) or (S3);
- (3) an expression vector (IV), or a recombinant live microorganism comprising (III);
- (4) a host cell (V) comprising (IV), or a subcellular fraction or membrane of the host cell expressing (I);
- (5) production of (I) comprising culturing (V) and recovering the produced polypeptide;
- (6) expressing (III) involves transforming (V) with (IV) which contains any one of the polynucleotides given above and culturing (V) under suitable conditions to express the polynucleotides;
 - (7) a vaccine composition which comprises (I) or (II);
 - (8) a vaccine composition which comprises (III);
 - (9) an antibody (Ab) immunospecific for (I) or (II); and
- (10) therapeutic compositions comprising an antibody directed against (I) useful in treating humans with Moraxella catarrhalis.

ACTIVITY - Anti-inflammatory; auditory; antibacterial.

MECHANISM OF ACTION - Gene therapy; vaccine. Details of test
are given but no results are stated.

USE - (I), (II) and (III) are useful for preparing a medicament useful for generating an immune response in an animal. (I) is also useful as diagnostic reagent for Moraxella catarrhalis which involves identifying (I) or an antibody against (I) present within the biological sample from an animal suspected of having such an infection (claimed). The polynucleotides may be used as hybridization probes for RNA, cDNA and genomic DNA to isolate full-length cDNAs and genomic clones encoding BASB113 and to isolate cDNA and genomic clones of other genes that have high sequence identity to BASB113 gene. The polynucleotides and polypeptides are used as research reagents and materials for discovery of treatments of and diagnostics for human diseases. The polynucleotides derived from (S1) or (S3) is used as PCR (polymerase chain reaction) primers. The polynucleotide sequences can be used in the discovery and development of antibacterial compounds. The encoded protein can be used as target for the screening of antibacterial drugs. Additionally, the polynucleotide sequences encoding the amino terminal regions of the encoded protein or Shine-Dalgarno or other translation facilitating sequences of the respective mRNA can be used to construct antisense sequences to control the expression of the coding sequence of interest. The polypeptides and polynucleotides are used to block the initial physical interaction between a gram negative and/or gram positive bacteria to mammalian, host thus preventing the sequelae of infection. The polynucleotides encoding certain non-variable regions of bacterial cell surface protein are used in polynucleotide constructs which are useful for genetic immunization experiments in animal models of infection with Moraxella catarrhalis to identify protein groups able to provoke a

prophylactic or therapeutic immune response. The vaccine comprising (I), (II) or (III) is useful for treating Moraxella catarrhalis infections such as sinusitis, nosocomial infections, otitis media and pneumonia. (II) is also used for therapeutic or prophylactic purposes especially genetic immunization.

Dwg.0/3

L26 ANSWER 16 OF 20 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2001-112457 [12] WPIDS

DOC. NO. NON-CPI: N2001-082525 DOC. NO. CPI: C2001-033486

TITLE: Novel BASB112 polypeptides of Moraxella

catarrhalis, useful as a vaccine for treating

Moraxella catarrhalis infections.

DERWENT CLASS: B04 D16 S03 INVENTOR(S): THONNARD, J

PATENT ASSIGNEE(S): (SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT: 95

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001000835 A1 20010104 (200112)* EN 81

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG

KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN

YU ZA ZW

AU 2000061519 A 20010131 (200124)

EP 1196591 A1 20020417 (200233) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	API	PLICATION	DATE
WO 2001000835 AU 2000061519 EP 1196591		AU EP	2000-EP5849 2000-61519 2000-947873 2000-EP5849	20000623 20000623 20000623 20000623

FILING DETAILS:

PATENT NO	KIND	PA	ATENT NO
AU 200006151 EP 1196591	9 A Based Al Based		200100835

PRIORITY APPLN. INFO: GB 1999-14870 19990625

AN 2001-112457 [12] WPIDS

AB WO 200100835 A UPAB: 20010302

NOVELTY - Isolated BASB112 polypeptides (I) of Moraxella catarrhalis, are new. The BASB112 polypeptide has the 122 (P1) or another 122 (P2) amino acid sequence defined in the specification.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for

the following:

- (1) an isolated polypeptide (Ia) comprising an amino acid sequence which is at least 85%, preferably 95%, most preferably 100%, identical to the sequence, over its entire length, of P1 or P2;
- (2) an immunogenic fragment (Ib) of (I) or (Ia), where the activity of the fragment is substantially the same as P1 or P2;
 - (3) an isolated polynucleotide (II) encoding (I), (Ia) or (Ib);
- (4) an isolated polynucleotide (IIa) comprising a sequence encoding (Ia) or its complementary sequence
- (5) an isolated polynucleotide (IIb) comprising a sequence having at least 85%, preferably 95%, most preferably 100% identity to a sequence encoding Pl or P2 over the entire coding region, or a nucleotide sequence complementary to the isolated polynucleotide;
- (6) an isolated polynucleotide (IIc) comprising a sequence having at least 85%, preferably 95%, most preferably 100% identical to the 369 (N1) or 366 (N2) nucleotides fully defined in the specification, or its complement;
- (7) an isolated polynucleotide (IId) comprising a sequence encoding P1 or P2, obtainable by screening an appropriate library under stringent hybridization conditions with labeled probe having the sequence of N1 or N2;
- (8) an expression vector (III) of a recombinant live microorganism, comprising (II), (IIa), (IIb), (IIc) or (IId);
- (9) a host cell (IV) comprising (III), or a subcellular fraction or membrane of (IV) expressing (Ia);
- (10) a process for producing (I), (Ia) or (Ib) comprising culturing (IV)
- (11) a process for expressing (II), (IIa), (IIb), (IIc) or (IId), comprising transforming (IV) with (III) and culturing transformed (IV) under conditions sufficient for its expression;
- (12) a vaccine composition (V) comprising (I), (Ia) or (Ib), or (II), (IIa), (IIb), (IIc) or (IId);
- (13) an antibody (Ab1) immunospecific for (I), (Ia) or (Ib); and
- (14) a method for diagnosing Moraxella catarrhalis infection, by identifying (I)-(Ib) or Ab1 present within a biological sample from an animal suspected of having such an infection.

ACTIVITY - Antibacterial.

MECHANISM OF ACTION - Vaccine.

Groups of mice are immunized with BASB112 vaccine. After the booster, the mice were challenged by instillation of bacterial suspension into the nostril under anaesthesia. Mice were killed between 30 minutes and 24 hours after challenge and the lungs were removed aseptically and homogenized individually. The log 10 weighted mean number of colony forming units (CFU)/lung was determined by counting the colonies grown on agar plates after plating of dilutions of the homogenate. The arithmetic mean of the log 10 weighted mean number of CFU/lung and the standard deviations were calculated for each group. Results were not given in the specification.

USE - The vaccine is useful for preparing a medicament for use in generating immune response in an animal (claimed). Abl is useful for treating humans suffering from Moraxella catarrhalis disease (claimed).

Polynucleotides encoding the BASB112 polypeptides have utility in diagnosis of the stage and type of infection, and also for therapeutic or prophylactic purposes, in particular genetic immunization. Dwg.0/3

L26 ANSWER 17 OF 20 HCAPLUS COPYRIGHT 2002 ACS DUPLICATE 2

ACCESSION NUMBER: 2000:133833 HCAPLUS

DOCUMENT NUMBER: 132:176650

TITLE: Cloning of BASB023 antigen from Moraxella

catarrhalis

INVENTOR(S):
Thonnard, Joelle

PATENT ASSIGNEE(S): Smithkline Beecham Biologicals S.A., Belg.

SOURCE: PCT Int. Appl., 99 pp.

CODEN: PIXXD2
Patent

DOCUMENT TYPE:

LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	PAT	ENT I	.00		KII	ND I	DATE				AP	PLIC	CATIO	ON NO	ο.	DATE		
	WO	20000	00969	94	A:	1 :	2000	0224			WO	199	99-EI	P582	8	1999	0811	
		W:	ΑE,	AL,	AM,	AT,	AU,	AZ,	BA,	BE	3,	BG,	BR,	BY,	CA,	CH,	CN,	CR,
																GM,		
			ID,	IL,	IN,	IS,	JP,	KE,	KG,	K	?,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,
			LU,	LV,	MD,	MG,	MK,	MN,	MW,	MΧ	ζ,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,
			SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TF	₹,	TT,	UA,	ŪĠ,	US,	UZ,	VN,	YU,
			ZA,	ZW,	AM,	AZ,	BY,	KG,	KZ,	ME	Ο,	RU,	ΤJ,	TM				•
		RW:	GH,	GM,	ΚE,	LS,	MW,	SD,	SL,	SZ	Ζ,	UG,	ZW,	ΑT,	BE,	CH,	CY,	DE,
			DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	ΙΊ	Γ,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,
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	EΡ	1105																
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GE	3,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,
			PT,	IE,	SI,	LT,	LV,	FI,										
PRIOR	RITY	APP	LN.	INFO	. :									_	-	1998		
										WO	19	99-E	EP582	28	W	19990	0811	

AB The invention provides BASB023 polypeptides and polynucleotides encoding BASB023 polypeptides from Moraxella catarrhalis (also named Branhamella catarrhalis) and methods for producing such polypeptides by recombinant techniques. BASB023 antigen is related by amino acid sequence homol. to Legionella adelaidensis macrophage infectivity potentiator polypeptide. Since Moraxella catarrhalis is responsible for several pathologies, the main ones being otitis media in infants

and children and pneumonia in elderlies, the invention provides diagnostic, prophylactic and therapeutic uses for Moraxella

infection.

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L26 ANSWER 18 OF 20 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2001-025166 [03] WPIDS

DOC. NO. NON-CPI: N2001-019583 DOC. NO. CPI: C2001-007779

TITLE: New BASB103-108 polypeptides isolated from Moraxella catarrhalis bacterium, useful for

diagnosing and producing vaccines against bacterial

infections such as otitis media and pneumonia.

DERWENT CLASS: B04 D16 S03

INVENTOR(S):

THONNARD, J

PATENT ASSIGNEE(S):

(SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT:

94

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2000071724 A2 20001130 (200103)* EN 79.

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000045673 A 20001212 (200115)

EP 1185658 A2 20020313 (200225) EN

R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI

APPLICATION DETAILS:

PATENT NO K	IND	APPLICATION	DATE
WO 2000071724 AU 2000045673 EP 1185658	• • • • • • • • • • • • • • • • • • • •	WO 2000-EP4618 AU 2000-45673 EP 2000-927226 WO 2000-EP4618	20000518 20000518 20000518 20000518

FILING DETAILS:

PATENT	NO F	KIND				TENT NO
AU 200	0045673	3 A	Based	on	WO	200071724
EP 118	5658	Α2	Based	on	WO	200071724

PRIORITY APPLN. INFO: GB 1999-13354 19990608; GB 1999-12038 19990524; GB 1999-12040 19990524; GB 1999-12674 19990601; GB 1999-12705 19990601; GB 1999-12838 19990602

AN 2001-025166 [03] WPIDS

AB WO 200071724 A UPAB: 20010116

NOVELTY - An isolated polypeptide (I) comprising an amino acid sequence which is at least 85% identical to the Moraxella catarrhalis BASB103-BASB108 polypeptides fully defined sequence of 252 (S2), 650 (S4), 405 (S6), 410 (S8), 818 (S10) or 913 (S12) amino acids as given in the specification, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

(1) an immunogenic fragment (II) of (I) which has the same immunogenic activity as (I);

(2) an isolated polynucleotide (III), or its complementary nucleotide sequence comprising a nucleotide sequence:

(a) encoding (I);

(b) that is 85% identical over the entire sequence which

encodes (S2), (S4), (S6), (S8), (S10) or (S12); (c) that is 85% identical to a fully defined nucleotide sequence of 759 (S1), 1953 (S3), 1218 (S5), 1233 (S7), 2457 (S9) or

- 2742 (S11) base pairs as given in the specification; and
- (d) comprising a nucleotide sequencing encoding (I) obtainable by screening an appropriate library under stringent hybridization conditions with a labeled probe having the sequence of (S1), (S3), (S5), (S7), (S9) or (S11);
- (3) an expression vector (IV) or a recombinant live microorganism comprising (III);
- (4) a host cell (V) comprising (IV), or a subcellular fraction or membrane of the host cell expressing (I);
 - (5) preparation of (I);
- (6) expressing (III) involves transforming (V) with (IV) and culturing (V) under suitable conditions to express the polynucleotides;
 - (7) a vaccine composition which comprises (I), (II) or (III);
 - (8) an antibody (Ab) immunospecific for (I) or (II); and
- (9) therapeutic compositions comprising an Ab directed against
- ACTIVITY Anti-inflammatory; auditory. No supporting data given.

MECHANISM OF ACTION - Gene therapy; vaccine.

USE - The therapeutic composition comprising (I), an immunogenic fragment (II) of (I) or a polynucleotide (III) encoding (I) is useful for the preparation of a medicament for generating an immune response in an animal. (I) is also useful as a diagnostic reagent for Moraxella catarrhalis which involves identifying (I) or an antibody against (I) present within the biological sample from an animal suspected of having such an infection (claimed). The polynucleotides may be used as hybridization probes for RNA, cDNA and genomic DNA to isolate full-length cDNAs and genomic clones encoding BASB103-108 and to isolate cDNA and genomic clones of other genes that have high sequence identity to BASB103-108 gene. The polynucleotides and polypeptides are used as research reagents and materials for discovery of treatments of and diagnostics for human diseases. The polynucleotides derived from (S1), (S3), (S5), (S7), (S9) or (S11) are used as polymerase chain reaction (PCR) primers. The polynucleotide sequences can be used in the discovery and development of antibacterial compounds. The encoded protein can be used as target for the screening of antibacterial drugs. Additionally, the polynucleotide sequences encoding the amino terminal regions of the encoded protein or Shine-Dalgarno or other translation facilitating sequences of the respective mRNA can be used to construct antisense sequences to control the expression of the coding sequence of interest. The polypeptides and polynucleotides are used to block the initial physical interaction between a gram negative and/or gram positive bacteria to mammalian host thus preventing the sequelae of infection. The polynucleotides encoding certain non-variable regions of bacterial cell surface protein are used in polynucleotide constructs which are useful for genetic immunization experiments in animal models of infection with M.catarrhalis to identify protein groups able to provoke a prophylactic or therapeutic immune response. The vaccine comprising (I), (II) or (III) is useful for treating Moraxella catarrhalis infections such as sinusitis, nosocomial infections, otitis media and pneumonia. (II) is also used for therapeutic or prophylactic purposes especially genetic immunization. Dwg.0/0

ACCESSION NUMBER: 2000-062301 [05] WPIDS DOC. NO. NON-CPI: N2000-048799

DOC. NO. CPI: C2000-017245

TITLE: Novel peptides useful as vaccines for Moraxella

infections such as otitis media, pneumonia,

sinusitis etc.,.

DERWENT CLASS: B04 D16 S03

INVENTOR(S): THOHNARD, J; THONNARD, J

PATENT ASSIGNEE(S): (SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT: 87

PATENT INFORMATION:

PAT	ENT	NO	F	KIND) D2	ATE		W	EEK		1	ĹΑ	PC	3							
WO	9958	3684	 !	A2	19	999:	1118	3 (2	2000	005)	* 1	EN	113	3							
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		MW	NL	OA	PT	SD	SE	SL	SZ	UG	zw										
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•				GD																	
		LR	LS	LT	LU	LV	MD	MG	MK	MN	MW	MX	NO	NZ	\mathbf{br} .	PΤ	RO	RU	SD	SE	SG
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EΡ	1078																			•	
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NO	2000	2005	697	7 A.	20	0010)11() (2	200:	115)									•		
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	991									225)											
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APPLICATION DETAILS:

PATENT NO KIND	APPLICATION	DATE
WO 9958684 A2	WO 1999-EP3257	19990507
AU 9941421 . A	AU 1999-41421	19990507
EP 1078064 A2	EP 1999-924948	19990507
	WO 1999-EP3257	19990507
NO 2000005697 A	WO 1999-EP3257	19990507
	NO 2000-5697	20001110
CZ 2000004203 A3	WO 1999-EP3257	19990507
·	CZ 2000-4203	19990507
AU 737196 B	AU 1999-41421	19990507
KR 2001043573 A	KR 2000-712705	20001113
CN 1309706 A	CN 1999-808554	19990507
HU 2001002853 A2	WO 1999-EP3257	19990507
	HU 2001-2853	19990507
ZA 2000006522 A	ZA 2000-6522	20001110
BR 9911773 A	BR 1999-11773	19990507
	WO 1999-EP3257	19990507
MX 2000011140 A1	MX 2000-11140	20001113
JP 2002514425 W	WO 1999-EP3257	19990507
	JP 2000-548475	19990507

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FILING DETAILS:	
PATENT NO KIND	PATENT NO
AU 9941421 A Based on	
EP 1078064 A2 Based on CZ 2000004203 A3 Based on	
AU 737196 B Previous Publ.	
. Based on	WO 9958684
HU 2001002853 A2 Based on	
BR 9911773 A Based on	
JP 2002514425 W Based on	WO 9958684
PRIORITY APPLN. INFO: GB 1998-10285	19980513
AN 2000-062301 [05] WPIDS	
AB WO 9958684 A UPAB: 20000128	
NOVELTY - An isolated polypeptide	with Moraxella catarrhalis
BASB020 polypeptide (1),(11),(111 (aa) as given in the specification),(IV) sequence of 280 amino acids
MC2931, MC2912, MC2913 and MC2969	is new
DETAILED DESCRIPTION - INDEP	PENDENT CLAIMS are also included for
the following:	
	(V), comprising an aa sequence
which has 85% identity to the aa	sequence of (I), (II), (III) or (IV);
(2) an immunogenic fragment	(VI), of (I),(II),(III),(IV) or
(V) which has the same immunogeni	c activity as (I),(II),(III) or
(IV);	A CONTROL OF THE CONT
	de (VII), comprising a nucleotide
sequence encoding (I).(II).(III)	or (IV):

leotide sequence encoding (I), (II), (III) or (IV);

- (4) an isolated polynucleotide (VII), or its complementary nucleotide sequence comprising a nucleotide sequence:
- (a) encoding a polypeptide that has 85% identity over the entire length of (I),(II),(III) or (IV);
- (b) that has 85% identity over the entire length of the nucleotide sequence coding region which encodes (I), (II), (III) or
- (c) which has 85% identity over the entire length of a fully defined nucleotide sequence of 843 base pairs (1,2,3,4) as given in the specification;
- (5) an expression vector (IX), or a recombinant live microorganism comprising (VII) or (VIII);
- (6) a host cell (X), or a membrane comprising (IX) which expresses (V);
 - (7) preparation of (I), (II), (III) or (IV);
- (8) expression of (VII) or (VIII) which comprises transforming (X) with (IX) which contains any one of the polynucleotides given above and culturing (X) under suitable conditions to express the polynucleotides;
- (9) a vaccine composition which comprises (I), (II), (III) or (IV) or (V);
 - (10) a vaccine composition which comprises (VII) or (VIII);
- (11) an antibody (Ab) immunospecific for (I), (II), (III), (IV), (V) or (VI); and
- (12) diagnosing a Moraxella infection by identifying (I),(II),(III), (IV),(V) or (VI) or an Ab produced against them, present in a biological sample obtained from an animal suspected of having such infection.

ACTIVITY - Anti-inflammatory; auditory.

MECHANISM OF ACTION - Vaccine. The efficacy of BASB020 vaccine was analyzed by enhancement of lung clearance of M.catarrhalis in mice. Groups of 6 BALB/c mice were immunized subcutaneously with 100 mu l of vaccine corresponding to a 10 mu l dose and were boosted 2 weeks later. One week after the booster, the mice were challenged by instillation of 50 mu l of bacterial suspension into the left nostril under anesthesia and 0.8 mg ketamine. Mice were killed 4 hours after challenge and the lungs are removed aseptically a homogenized individually. The log 10 weighted mean number of CFU/lung is determined by counting the colonies grown on Mueller-Hinton agar plates after plating of 20 mu l of 5 serial dilutions of the homogenate. BASB020 vaccine induced significant lung clearance as compared to the control (0.62 log difference).

USE - The polynucleotides may be used as hybridization probes for RNA, cDNA and genomic DNA to isolate full-length cDNAs and genomic clones encoding BASB020 and to isolate cDNA and genomic clones of other genes that have high sequence identity to BASB020 gene. The polynucleotides and polypeptides are used as research reagents and materials for discovery of treatments of and diagnostics for human diseases. The polynucleotides derived from (1,2,3,4) are used for PCR to determine whether or not the identified polynucleotides are transcribed in bacteria in infective tissue and so are helpful in the diagnosis of the stage and type of infection, the pathogen has attained. Probes comprising BASB020 nucleotide sequence can be constructed to conduct efficient screening of genetic mutations, serotype, taxonomic classification or identification. Primers with 1-4 nucleotides removed from the 5' and/or 3' end are used for amplifying BASB020 DNA and/or RNA isolated from a sample derived from an individual. The polynucleotides are used as components of high density polynucleotide arrays or grids which are useful for diagnostic and prognostic purposes. The antibodies directed against (I),(II),(III),(IV) or (VII) are employed to isolate or to identify clones expressing (I),(II),(III),(IV) or (VII) or to purify them. The polynucleotide sequences can be used in the discovery and development of antibacterial compounds. The encoded protein, for expression can be used as target for the screening of antibacterial drugs. Additionally, the polynucleotide sequences encoding the amino terminal regions of the encoded protein or Shine-Delgarno or other translation facilitating sequences of the respective mRNA can be used to construct antisense sequences to control the expression of the coding sequence of interest. The polypeptides and polynucleotides are used to block the initial physical interaction between a gram negative and/or gram positive bacteria to the mammalian host. The polynucleotides encoding certain non-variable regions of bacterial cell surface protein are used in polynucleotide constructs which are useful for genetic immunization experiments in animal models of infection with M.catarrhalis to identify protein epitopes able to provoke a prophylactic or therapeutic immune response. The therapeutic composition comprising an immunologically effective amounts of a polypeptide, (I),(II),(III),(IV) or (V); or a polynucleotide, (VII) or (VIII) is useful in the preparation of a medicament for generating an immune response in an animal. A therapeutic composition comprising an Ab directed against one or two useful for treating humans with M.catarrhalis diseases (claimed) such as sinusitis, otitis media and nosocomial infections. Dwq.0/8

L26 ANSWER 20 OF 20 WPIDS (C) 2002 THOMSON DERWENT

ACCESSION NUMBER: 2000-039107 [03] WP:

DOC. NO. NON-CPI: N2000-029453 DOC. NO. CPI: C2000-010168

TITLE: Novel BASB010 polynucleotides and polypeptides from

Moraxella catarrhalis used to prepare vaccines

against bacterial infections.

DERWENT CLASS: B04 D16 S03 INVENTOR(S): THONNARD, J

PATENT ASSIGNEE(S): (SMIK) SMITHKLINE BEECHAM BIOLOGICALS

COUNTRY COUNT: 87

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 9958682 A2 19991118 (200003)* EN 100 RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC

MW NL OA PT SD SE SL SZ UG ZW

W: AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES

FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG

SI SK SL TJ TM TR TT UA UG US UZ VN YU ZA ZW

AU 9942600 A 19991129 (200018)

EP 1078065 A2 20010228 (200113) EN

R: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 9958682 AU 9942600 EP 1078065	A2 A A2	WO 1999-EP3254 AU 1999-42600 EP 1999-950353 WO 1999-EP3254	19990507 19990507 19990507 19990507

FILING DETAILS:

PATENT NO	KIND	PATENT NO
AU 9942600	A Based on	WO 9958682
EP 1078065	A2 Based on	WO 9958682

PRIORITY APPLN: INFO: GB 1999-5308 19990308; GB 1998-10195

19980512

AN 2000-039107 [03] WPIDS

AB WO 9958682 A UPAB: 20000118

NOVELTY - Novel BASB010 polynucleotides and polypeptides from Moraxella catarrhalis are disclosed.

DETAILED DESCRIPTION - An isolated BASB010 polypeptide (I) is new, and comprises an amino acid sequence which has at least 85% or 95% identity to, or is, the 391 (Ia), 391 (Ib) or 391 (Ic) amino acid sequences given in the specification.

INDEPENDENT CLAIMS are also included for the following:

(1) An immunogenic fragment of (I) in which the immunogenic activity is substantially the same as (Ia), (Ib) or (Ic);

(2) An isolated polynucleotide encoding (I), or a complementary nucleotide;

- (3) An isolated polynucleotide (II) which comprises a sequence which has at least 85% or 95% identity to over the entire length, or is, the 1176 bp (IIa), 1176 bp (IIb) or 1176 bp (IIc) sequence given in the specification, or its complement;
- (4) An isolated polynucleotide encoding (Ia)-(Ic), obtainable by screening an appropriate library under stringent hybridization conditions with a labeled probe having the sequence of (IIa), (IIb), (IIc) or a fragment thereof;
- (5) An expression vector or recombinant live microorganism comprising (II), or the polynucleotides of (2) or (4);
- (6) A host cell comprising the expression vector of (5), or a subcellular fraction of that cell expressing (I);
- (7) A process for producing (I), comprising culturing a host cell under conditions sufficient for the production of the polypeptide, and recovering the polypeptide from the culture medium;
- (8) A process for expressing (II) or the polynucleotides of (2) or (4), comprising transforming a host cell with a vector comprising at least one of these polynucleotides, and culturing the cell under conditions sufficient for expression of the polynucleotide;
- (9) A vaccine composition comprising an effective amount of (I) and a pharmaceutically acceptable carrier;
- (10) A vaccine composition comprising an effective amount of (II) or the polynucleotides of (2) or (4), and a pharmaceutically acceptable carrier;
- (11) An antibody immunospecific for (I), or the fragment of (1);
- (12) A method for diagnosing a M. catarrhalis infection, comprising identifying (I), or an antibody that is immunospecific for (I), present within a biological sample from an animal suspected of having such an infection;
- (13) Use of a composition comprising an immunologically effective amount of (I) or (II) or the polynucleotides of (2) or (4) in the preparation of a medicament for use in generating an immune response in an animal; and
- (14) A therapeutic composition useful in treating humans with M. catarrhalis, comprising at least one antibody directed against (I) and a pharmaceutically acceptable carrier.

ACTIVITY - Anti-bacterial, immunostimulant.

MECHANISM OF ACTION - Vaccine.

USE - The polynucleotides and polypeptides may be employed as research reagents and material for the discovery of treatments and diagnostics for diseases, particularly human diseases. They can be used for diagnosis of disease, staging of disease, or determining response of an infectious organism to drugs. The polynucleotides may be used as a source for hybridization probes, and for screening of genetic mutations, serotype, organism or strain identification, identification of mutations in BASB013 sequences, and as components of arrays which are useful for diagnostic and prognostic purposes. The polypeptides can be used to produce antibodies. The polypeptides can also be used in vaccine formulations, and to identify agonists and antagonists. The polypeptides, antibodies, agonists and antagonists (which are bacteristatic) are used for the treatment and prevention of diseases such as otitis media in infants and children, pneumonia in the elderly, sinusitis, nosocomial infections and invasive diseases, chronic otitis media with hearing loss, fluid accumulation in middle ear, auditive nerve damage, delayed speech learning, infection of the upper respiratory tract and inflammation of the middle ear. They are particularly used to diagnose and treat

M. catarrhalis infections. The polypeptides, agonists and antagonists are also used for screening of antibacterial drugs.

ADVANTAGE - The frequency of Moraxella catarrhalis infections has risen dramatically, and it is no longer common to isolate M. catarrhalis strains that are resistant to standard antibiotics. The BASB010 products of the invention can be used screen for new antibacterial compounds that may target these resistant bacteria. Dwg.0/4

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